

# Latest Cretaceous megaraptorid theropod dinosaur sheds light on megaraptoran evolution and palaeobiology

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Recent fossil discoveries have cast considerable light on the palaeobiology of Megaraptora, a group of large-clawed carnivorous theropod dinosaurs known from Cretaceous deposits in Asia, Australia, and especially South America. Nevertheless, many important aspects of megaraptoran morphology and evolution remain poorly understood, due in large part to the fragmentary nature of most fossils of these theropods and the scarcity of anatomically overlapping skeletal elements among the known taxa. Here we report a previously unknown megaraptoran genus and species represented by a partially articulated partial skeleton recovered from an uppermost Cretaceous stratum of the Lago Colhué Huapi Formation of south-central Chubut Province in central Patagonia, Argentina. Pertaining to the derived megaraptoran subclade Megaraptoridae, the taxon is among the most completely represented and latest-surviving megaraptorans. Its stratigraphic occurrence indicates that these dinosaurs likely persisted to the Cretaceous/Palaeogene boundary; moreover, the preservation of a crocodyliform humerus between the dentaries of the new theropod may provide information on megaraptoran dietary preferences and feeding strategies. Megaraptorids appear to have been the apex predators in central and southern Patagonian palaeoecosystems approaching the end of the Cretaceous, in contrast to more northerly areas of South America where these niches were occupied by other non-avian theropod groups.

Among the most mysterious of the many predominantly Gondwanan Cretaceous non-avian theropod dinosaur groups is Megaraptora, a tetanuran clade characterized by (at least in derived members of the group) elongate skulls with low-crowned teeth, extensively pneumatized bones, and, most strikingly, powerfully developed forelimbs

equipped with hypertrophied unguals on the first and second digits<sup>1–9</sup>. Although the stratigraphically oldest and earliest diverging representatives of this clade come from Asia<sup>2,10–13</sup> and Australia<sup>14–19</sup>, megaraptorans are particularly abundant in South America, especially in the Late Cretaceous of Patagonia (e.g., refs. 1,3,4,6–8,20–32; see also

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Supplementary Notes 1 and 2). The evolutionary relationships of Megaraptora within Tetanurae—as either carcharodontosaurian allosauroids or early branching coelurosaurids, potentially basal tyrannosauroids—are controversial (see refs. 6,8,21,25,33–36, among others), due in large part to the preponderance of fragmentary, highly incomplete fossils of the clade and the scarcity of anatomically overlapping skeletal elements among the various megaraptoran taxa.

Here we report a new megaraptoran genus and species based on a well-preserved, partly articulated partial skeleton that includes cranial, postcranial axial, and appendicular elements, recovered from the Upper Cretaceous (Coniacian–Maastrichtian) Lago Colhué Huapi Formation of south-central Chubut Province, central Patagonia, Argentina (see Supplementary Note 3 for additional locality and stratigraphic data). Specifically, the new taxon, a member of the derived megaraptoran subclade Megaraptoridae, comes from a stratigraphic horizon only a few meters below of the top of this formation, establishing it as latest Cretaceous (Maastrichtian, probably late Maastrichtian) in age, and therefore one of the geologically youngest (if not the youngest) megaraptorans yet discovered. Moreover, because this taxon is also one of the most completely represented megaraptorans, it adds significantly to our understanding of the morphology, phylogenetic relationships, and palaeobiology of this enigmatic large-clawed theropod group. Furthermore, the unusually close association of a crocodyliform humerus with the type specimen of this theropod may provide information on the dietary preferences and feeding strategies of Megaraptoridae.

## Results and discussion

### Systematic palaeontology

Saurischia Seeley, 1887<sup>37</sup>

Theropoda Marsh, 1881<sup>38</sup>

Tetanurae Gauthier, 1986<sup>39</sup>

Megaraptora Benson, Carrano, and Brusatte, 2010<sup>33</sup>

Megaraptoridae Novas, Agnolín, Ezcurra, Porfiri, and Canale, 2013<sup>34</sup>

*Joaquinraptor casali* gen. et sp. nov.

**Etymology.** Joaquín, in tribute to the son of the first author (L.M.I.) and the informal name given to the locality when the skeleton of the taxon was discovered (Valle Joaquín); Latin, raptor, thief. Specific epithet *casali* in recognition of Dr. Gabriel Andrés Casal for his many contributions to knowledge of the Cretaceous palaeontology and geology of central Patagonia (including the formal recognition and naming of the formation from which this megaraptorid was recovered).

**Holotype.** UNPSJB-PV 1112, a partially articulated partial skeleton of a single individual that includes the mostly disarticulated partial skull (right maxilla, skull roof and braincase, probable right postorbital, right quadrate, both dentaries, in situ and isolated teeth), complete or nearly complete elements of the postcranial axial (atlantal intercentrum, three caudal vertebrae, dorsal ribs, other rib and probable gastralia fragments, haemal arch) and appendicular (left scapulo-coracoid, humerus, radius, and ulna, right manual ungual II, left femur, much of the right tibia, the distal end of the probable right pedal phalanx III-3, the right pedal ungual III, and the possible partial left pedal phalanx IV-1) skeletons, and numerous indeterminate fragments.

**Locality and horizon.** Valle Joaquín, headwaters of the Río Chico, east of the southeastern shore of Lago Colhué Huapi, Chubut Province, central Patagonia, Argentina. Upper (Maastrichtian, probably upper Maastrichtian) stratum of the Upper Cretaceous Lago Colhué Huapi Formation (see below; also Fig. 1 and Supplementary Figs. 1 and 2).

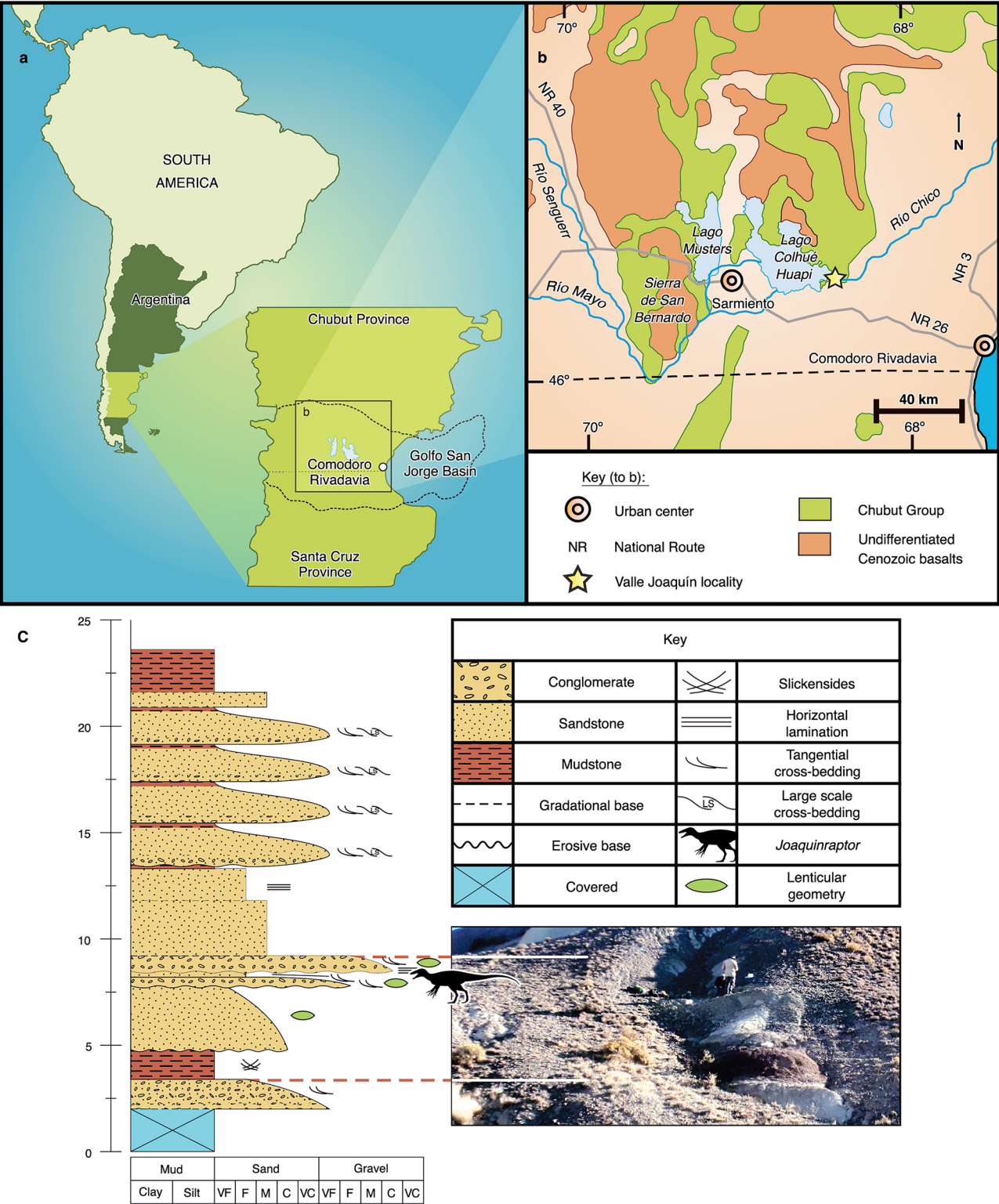
**Diagnosis.** The holotype of *Joaquinraptor casali* is considered to represent a sexually but possibly not somatically mature individual based on the fusion of braincase sutures and the results of our

osteohistological analysis, which reveal a reduction in the distance between growth marks in the outer bony cortex of limb bones (see below). *Joaquinraptor* is regarded as a member of Megaraptoridae based on its possession of the following suite of synapomorphies of the clade: (1) reduced anteriormost dentary alveolus<sup>35</sup>; (2) absence of mesial denticles from tooth crowns<sup>34,35,40,41</sup>; (3) olecranon process of ulna mediolaterally compressed, blade-like, and subtriangular in lateral view; and (4) ulnar shaft with well-developed, proximodistally oriented lateral crest or tuberosity<sup>21,34–36</sup>. *Joaquinraptor* is diagnosed by the following autapomorphies (see also Supplementary Note 4 and Supplementary Figs. 3–7): (1) parietals strongly compressed transversely, narrower than frontals; (2) anterior margin of supratemporal fossa with sigmoid contour; (3) braincase lacking median septum in pneumatic recess (probably, but see below); (4) rugose atlantal intercentrum with well-marked lips and rectangular ventral contour; (5) middle or posterior caudal vertebral centra with well-marked elliptical lateral fossae that occupy more than half the lateral surfaces; (6) humerus with subrectangular deltopectoral crest; ulna with (7) proximodistally enlarged olecranon process, (8) vertical ridge on medial surface of proximal end, (9) nearly straight shaft (but see below), and (10) medially oriented, anteroposteriorly compressed distal end; and (11) distal end of radius crescent-shaped. Differs from the only other named Maastrichtian (probably early Maastrichtian and therefore stratigraphically older; see refs. 42–44) megaraptorid species *Maip macrothorax* in having a coracoid with a subglenoid ridge, posteroventral fossa, and hooked posteroventral process. The preserved dorsal ribs of *Joaquinraptor* and *Maip* also exhibit morphological differences, though these could potentially be due to non-equivalence in serial position rather than taxonomic distinction.

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**Age.** Geological studies of the uppermost part of the Lago Colhué Huapi Formation, particularly those of palynomorphs, support a late Maastrichtian age for these strata, probably close to the Cretaceous/Palaeogene boundary<sup>45,46</sup>. The palaeoclimate of central Patagonia during deposition of this part of the formation is suggested to be warm with at least seasonal rainfall<sup>45</sup>. On the other hand, palaeoclimatic studies of the middle (Santonian –?lower Maastrichtian) section of the Lago Colhué Huapi Formation indicate a seasonally dry or semiarid climate<sup>47</sup>. Recent analysis of clays from the *Joaquinraptor* quarry indicates a composition similar to that of the top of the formation (see Supplementary Note 3). Therefore, this megaraptorid was found at a stratigraphic level that is more related to those of the top of this unit than those of the middle section, supporting a Maastrichtian (and probably late Maastrichtian) age for this specimen.

The oldest megaraptorans are *Phuwiangvenator yaemniyomi* and *Vayuraptor nongbualamphuensis* from Thailand<sup>11–13</sup>, both probably late Valanginian–early Hauterivian in age<sup>48</sup>. The youngest records of the clade are from the Campanian–Maastrichtian of Argentina, Chile, and possibly Brazil<sup>8,22,23,49–53</sup>. Nevertheless, these latter megaraptorans probably date to the Campanian–early Maastrichtian interval<sup>8,42–44,52–54</sup> and are thus similar in age to the fragmentary megaraptorid fossils (mostly manual unguals) recovered from the middle section of the Lago Colhué Huapi Formation<sup>7,31,55</sup>. Therefore, *Joaquinraptor* seemingly constitutes the stratigraphically youngest representative of Megaraptora yet discovered anywhere in



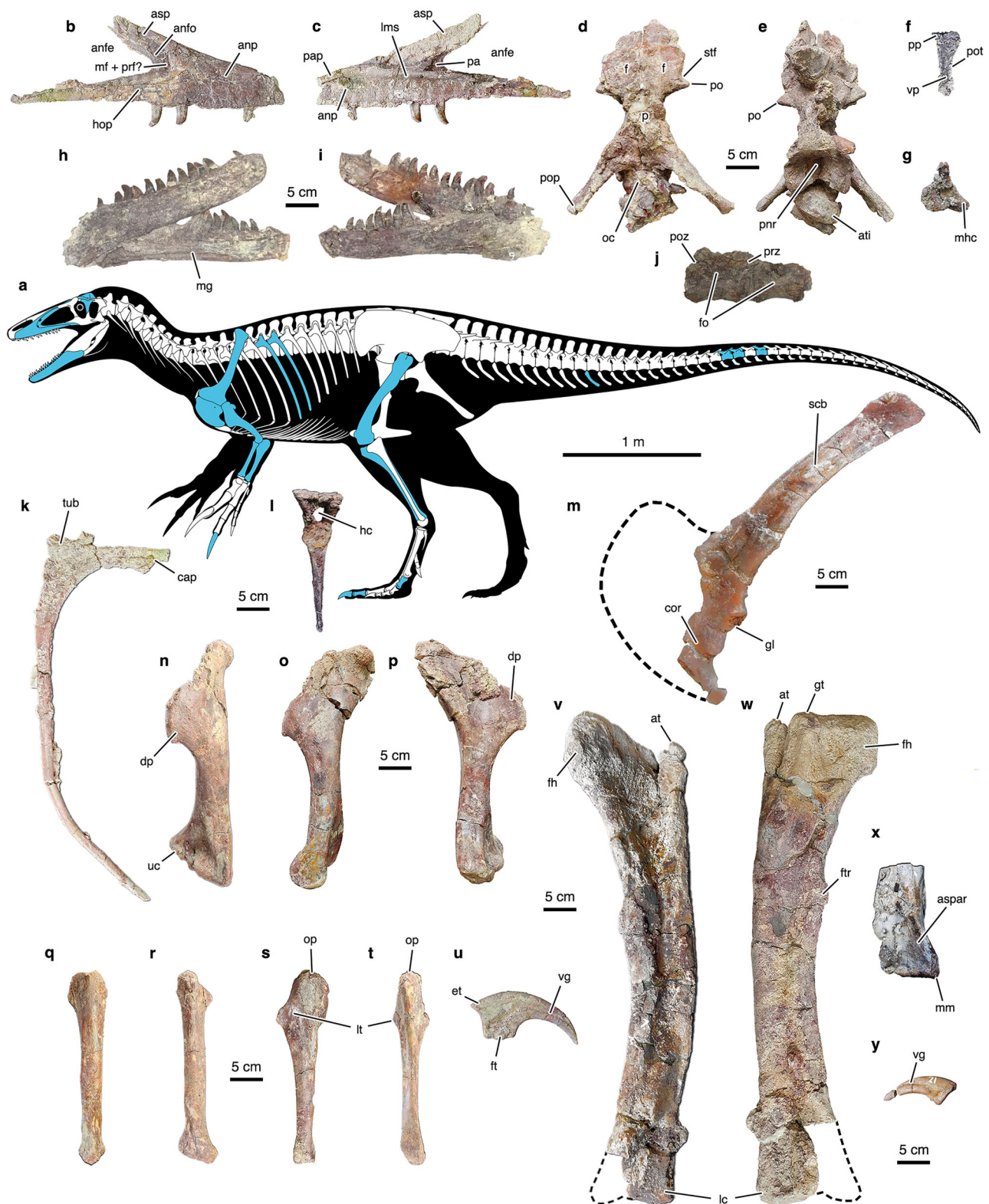
**Fig. 1 | Geographic and geological context of *Joaquinraptor casali* gen. et sp. nov.** **a** Location of the study area in south-central Chubut Province, central Patagonia, Argentina. **b** Simplified geologic map showing the informally named Valle Joaquín locality in the Upper Cretaceous Lago Colhué Huapi Formation (part

of the Chubut Group) that yielded *Joaquinraptor casali* gen. et sp. nov. **c** Stratigraphic column and photograph of Valle Joaquín showing the position of the horizon of the Lago Colhué Huapi Formation that yielded *Joaquinraptor*. Abbreviations: C coarse, F fine, M medium, VC very coarse, VF very fine.

the world. Interestingly, what are probably both the oldest<sup>7</sup> and youngest (*Joaquinraptor*) definitive South American megaraptorids known at present come from strata of the Golfo San Jorge Basin of central Patagonia.

**Osteological description.** The megaraptoran fossil record is generally depauperate; therefore, not many specimens within the group preserve overlapping skeletal elements. In this context, *Joaquinraptor casali* is important because it is represented by a well-preserved partial





skeleton that includes cranial, postcranial axial, and appendicular elements (Fig. 2a and Tables 1, 2), allowing relatively extensive comparisons with other megaraptoran taxa (see Supplementary Notes 5 and 6 and Supplementary Figs. 8–18). This is the case for the subtriangular maxilla (Fig. 2b, c), which is otherwise preserved only in a juvenile specimen of *Megaraptor namunhuaiquii* (MUCPv 595; see ref. 6). In *Joaquinaraptor*, the anterior part of the antorbital fossa is perforated by a poorly delimited, fossa-like structure that may

represent the confluent maxillary fenestra and promaxillary foramen. The morphology and configuration of these probable pneumatic excavations closely resembles that in *Megaraptor*, supporting this condition as a synapomorphy of Megaraptora or one of its subclades instead of an autapomorphy of the latter genus (*contra*<sup>6</sup>). Similarly, in both these megaraptorids, the medial maxillary shelf bifurcates posteriorly, forming a deep, “horizontal V”-shaped, well-defined antrum (=posterior anteromaxillary fenestra; see ref. 56), which may represent

**Fig. 2 | Osteology of *Joaquinraptor casali* gen. et sp. nov.** **a** Skeletal reconstruction of *Joaquinraptor* in left lateral view with preserved elements in blue (some reversed from right side) (modified and updated from Lamanna et al.<sup>7</sup> [these authors' Fig. 1e], which was in turn modified by A. McAfee from an original illustration by T.K. Robinson). **b** Right maxilla in lateral and medial (**c**) views. Skull roof, braincase, and atlantal intercentrum in dorsal (**d**) and ventral (**e**) views. **f** Probable right postorbital in lateral view. **g** Right quadrate in anterior view. Right and left dentaries in lateral and medial views (right dentary in lateral view and left dentary in medial view in **h**; opposite in **i**). **j** Two articulated middle or posterior caudal vertebrae in right lateral view. **k** Dorsal rib in anterior view. **l** Anterior haemal arch in anterior view. **m** Left scapulocoracoid in lateral view. Left humerus in anterolateral (**n**), lateral (**o**), and medial (**p**) views. Left radius in anterior (**q**) and lateral (**r**) views. Left ulna in lateral (**s**) and posterior (**t**) views. **u** Right manual ungual II (=manual phalanx II-3) in lateral view. Left femur in anterior (**v**) and posterior (**w**) views. **x** Distal right tibia in anterior view. **y** Right pedal ungual III (=pedal

phalanx III-4) in medial view. Dashed lines indicate missing areas of scapulocoracoid and femur. Abbreviations: anfe antorbital fenestra, anfo antorbital fossa, anp anterior process, aspar articular surface for ascending process of astragalus, at anterior trochanter, ati atlantal intercentrum, cap capitulum, cor coracoid, dp deltopectoral crest, et extensor tubercle, f frontal, fh femoral head, fo lateral fossa, ft flexor tubercle, ftr fourth trochanter, gl glenoid fossa, gt greater trochanter, hc haemal canal, hop horizontal process, lc lateral condyle, lms longitudinal medial shelf, lt lateral tuberosity, mg Meckelian groove, mf maxillary fenestra, mhc medial hemicondyle, mm medial malleolus, oc occipital condyle, op olecranon process, p parietal, pa posterior antrum, pap palatal process, pnr pneumatic recess, po articular surface for postorbital, pop paroccipital process, pot postorbital tuberosity, poz postzygapophysis, pp posterior process, prf promaxillary foramen, prz prezygapophysis, scb scapular blade, stf supratemporal fenestra, tub tuberculum, uc ulnar condyle, vg vascular groove, vp ventral process.

**Table 1 | Measurements (mm) of cranial and postcranial axial skeletal elements of *Joaquinraptor casali* gen. et sp. nov. (UNPSJB-PV 1112)**

Maxilla (R)	
Anteroposterior length (straight line)	403
Maximum dorsoventral height (excluding teeth)	130
Dentary (R)	
Anteroposterior length (straight line)	330
Symphysis dorsoventral height	45
Maximum dorsoventral height (excluding teeth)	65
Skull roof and braincase* (dorsal view)	
Anteroposterior length (along lateral margin)	195
Minimum transverse width (excluding paroccipital process)	50
Maximum transverse width (including paroccipital process)	120
Postorbital* (R)	
Dorsoventral length	94
Minimum anteroposterior width	25
Maximum anteroposterior width	41
Quadrate* (R)	
Maximum mediolateral width	65
Preserved dorsoventral height	75
Middle or posterior caudal vertebrae (articulated)	
Anteroposterior length, centra	75/72
Dorsoventral height/transverse width, anterior articular surface	50/43
Dorsoventral height/transverse width, posterior articular surface	40/45
Posterior caudal vertebra (isolated)	
Anteroposterior length, centrum	72
Dorsoventral height/transverse width, anterior articular surface	50/40
Anterior haemal arch	
Dorsoventral height (anterior view)	180
Maximum transverse width (anterior view)	62
Midshaft anteroposterior width (lateral view)	20

L left, R right, \* incomplete element.

another megaraptoran synapomorphy. The maxilla of *Joaquinraptor* differs from that of *Megaraptor* in having a substantially taller and blunter anterior process (=anterior ramus), an anteroposteriorly narrower anterior part of the antorbital fossa, an acute anterior margin of the antorbital fenestra, and a dorsoventrally thicker ascending process, although whether these distinctions are of taxonomic significance rather than reflective of the differing ontogenetic stages of UNPSJB-PV 1112 and MUCPv 595 is unknown at present. Interestingly, in being low, elongate, and subtriangular, the maxilla of *Joaquinraptor* is also somewhat similar to that of the penecontemporaneous large-

**Table 2 | Measurements (mm) of appendicular skeletal elements of *Joaquinraptor casali* gen. et sp. nov. (UNPSJB-PV 1112)**

Scapulocoracoid (L)	
Total anteroposterior length (including coracoid)	470
Maximum mediolateral width (ventral view)	30
Coracoid anteroposterior length*	95
Humerus (L)	
Maximum proximodistal length	330
Posterior midshaft mediolateral width	45
Radius (L)	
Maximum proximodistal length	215
Midshaft mediolateral width	25
Ulna (L)	
Maximum proximodistal length	271
Midshaft mediolateral width	30
Manual ungual II (R)	
Maximum proximodistal length (straight line)	140
Articular facet dorsoventral height	55
Femur* (L)	
Proximodistal length	685
Shaft mediolateral width (anterior view, straight line, 4th trochanter)	107
Midshaft circumference (just distal to 4th trochanter)	275
Pedal ungual III (L)	
Maximum proximodistal length (straight line)	95
Articular facet dorsoventral height	40

L left, R right, \* incomplete element.

bodied unenlagiine theropod *Austroraptor cabazai* from northern Patagonia<sup>57</sup>. Nevertheless, the maxilla of the new megaraptorid differs from that of *Austroraptor* in, among other features, the degree of elongation of the antorbital region (which is pronounced in the unenlagiine), the shape of the antorbital fossa, and the number of alveoli (which is substantially higher in the unenlagiine). Although the surfaces of the maxillary interdental plates are slightly eroded in the *Joaquinraptor* holotype, they appear to be unfused, unlike those of *Fukuiraptor kitadaniensis*<sup>2</sup>. The interdental plates are triangular in shape and subequal in size, as in the juvenile *Megaraptor* (MUCPv 595). On the other hand, allosauroids more derived than *Sinraptor dongi* have larger interdental plates<sup>6</sup>.

The braincase (Fig. 2d, e) is anteroposteriorly longer and more transversely compressed than that of the reportedly subadult holotype of the megaraptorid *Murusraptor barrosaensis* (MCF-PVPH 411; see ref. 25)—which is, at present, the only other almost complete megaraptoran braincase known—and still more narrow and elongate than

the partial braincase of the aforementioned juvenile *Megaraptor* (MUCPv 595; see ref. 6). No sutures are visible (except for the interparietal suture, see immediately below), which accords with osteohistological evidence that indicates that the *Joaquinraptor* holotype was sexually and possibly somatically mature at the time of death (minimum age of 19 years; see below). Nevertheless, although it has possibly been exaggerated by potential taphonomic artifacts, a well-marked, non-interdigitated, open interparietal suture is evident, as is also the case in *Murusraptor*; this feature may therefore characterize Megaraptoridae<sup>27</sup>. The parietals are more transversely constricted than in *Murusraptor*, closely resembling the condition in tyrannosauroids<sup>36</sup>; this feature is therefore considered an autapomorphy of *Joaquinraptor* within Megaraptora (Supplementary Fig. 3). The frontals are subquadrangular in shape, as in other megaraptorans<sup>6,18,24,25</sup>. In lateral view, the posterior region of the ventrolaterally oriented frontals occupies a more dorsal position than the anterior portion, conferring a step-like morphology similar to that of *Murusraptor* (see ref. 27) and the probable megaraptorid MCF-PVPH 320 (see ref. 24); this feature may therefore represent another synapomorphy of Megaraptora or a clade therein. Anteriorly, there is a wide, anteriorly projected supratemporal fossa similar to that seen in *Megaraptor* and *Murusraptor*, although it differs from those of these other megaraptorids in having a sigmoid contour in dorsal view. Conversely, in allosauroids such as *Acrocanthosaurus atokensis*, *Allosaurus fragilis*, and *Carcharodontosaurus saharicus*, the supratemporal fossa is anteroposteriorly short<sup>36</sup> and its anterior margin exhibits a more ovoid aspect. The ventrolateral inclination of the frontals from the midline of the skull roof, forming a triangular cross-section, has been regarded as an autapomorphy of *Murusraptor* (see ref. 27), but may instead represent a synapomorphy of Megaraptoridae as it is also present in *Joaquinraptor*. Ventrally, the median septum that divides the basisphenoid recess that is described in *Megaraptor*<sup>6</sup> and *Murusraptor*<sup>27</sup> appears to be absent in *Joaquinraptor*, although this could potentially be an artifact of preservation. The more complete of the two basipterygoid processes is robust and hooks posteriorly in lateral view, unlike those of *Murusraptor* that are shorter and mostly ventrally oriented.

The bone tentatively interpreted as the right postorbital (Fig. 2f) is almost complete, missing only parts of the anterior and posterior processes, though it has probably also experienced some plastic deformation. It is anteroposteriorly and mediolaterally narrower than the postorbitals of *Aerosteon riocoloradensis*<sup>4,30</sup>, *Murusraptor*<sup>25</sup>, and *Orkoraptor burkei*<sup>22</sup>. Compared with these megaraptorids, the anterior process appears shorter and the ventral process straighter (i.e., the latter is not ‘cradle-shaped’). *Joaquinraptor* shares with *Aerosteon* and *Murusraptor* the presence of a small tuberosity on the anterior margin of the ventral process that is well exposed in lateral and anterior views (see ref. 30, these authors’ Fig. 4a, d, “rugose bump”). Furthermore, both *Joaquinraptor* and *Aerosteon* possess a well-marked foramen on the dorsal surface of the postorbital, situated relatively close to the frontal contact.

The condylar (ventral) portion of the right quadrate is preserved (Fig. 2g). In overall morphology, it resembles the quadrate of *Aerosteon* more than that of *Murusraptor*, which is generally more gracile. There is a well-marked, enclosed posterior pneumatic foramen (*sensu*<sup>58</sup>) on the posterior surface, as in other megaraptorids and many other coelurosaurs<sup>58</sup>. In *Joaquinraptor*, however, this foramen is placed on the lateral part of the quadrate adjacent to the quadratojugal contact, whereas in *Aerosteon* and *Murusraptor* it is more centrally placed.

An isolated maxillary tooth of *Joaquinraptor* (Supplementary Fig. 17) is proportionally apicobasally taller than those of *Murusraptor*. As in other megaraptorids, the tooth is labiolingually compressed, the distal margin of the crown is strongly concave in labial and lingual views, and the distal denticles are subrectangular to rectangular. The

mesial margin lacks a serrated carina, as in most other South American megaraptorids<sup>40</sup>.

The *Joaquinraptor* dentaries (Fig. 2h, i) are elongate, mediolaterally compressed, and shallow in lateral view, as in *Australovenator wintonensis* (AODF 604)<sup>59</sup>. Furthermore, both these megaraptorids and the early branching megaraptoran *Fukuiraptor*<sup>2</sup> share a reduced symphyseal facet and an anterior dentary margin that trends anterodorsally–posteroventrally. However, in overall morphology, the dentary of *Joaquinraptor* is more robust than that of *Australovenator*. Slightly pronounced primary and secondary neurovascular foramina are evident on the lateral surface. White et al.<sup>59</sup> suggested that the absence of these foramina may be autapomorphic of *Australovenator*, contrasting with the original description of this taxon (see ref. 5). Similar neurovascular openings were described and figured in dentary fragments of *Fukuiraptor*<sup>2</sup>; therefore, the presence of slightly marked neurovascular foramina that are not enclosed or delimited by grooves may represent a synapomorphy of Megaraptora or one of its subclades (e.g., Megaraptoridae). Both *Joaquinraptor* dentaries preserve 17 alveoli—two fewer than in *Australovenator*—although this could be a taphonomic artifact caused by erosion of the posteriormost portions of these bones. Unlike those of *Fukuiraptor* and *Australovenator*, the dentary interdental plates are unfused.

The atlantal intercentrum is articulated with the occipital condyle (Fig. 2e and Supplementary Fig. 3b), such that most of its anterior features cannot be described in detail; however, it is rugose and disc-shaped, differing from its more crescentic counterparts in the megaraptorans *Aerosteon*<sup>4,30</sup>, *Orkoraptor*<sup>22</sup>, and *Phuwiangvenator*<sup>11</sup>. Ventrally, the intercentrum exhibits well-marked rims and is rectangular in shape, unlike other megaraptorans in which this element is preserved.

The three preserved caudal vertebrae belong to the middle or posterior section of the tail, being substantially longer anteroposteriorly than tall dorsoventrally (see Table 1). The centra (Fig. 2j) are hourglass-shaped in ventral view, and their lateral surfaces are excavated by well-developed, elliptical fossae, as in many other definitive or possible megaraptorans (e.g.,<sup>4,7,22,30,49,50,60,61</sup>); however, the fossae seen in *Joaquinraptor* are proportionally very large, encompassing more than half the length of the lateral surfaces, a unique feature of this megaraptorid (Supplementary Fig. 4). Dorsal ribs (Fig. 2k) are pneumatized and generally resemble those of other megaraptorids, such as *Australovenator*<sup>5</sup>, *Maip macrothorax*<sup>8</sup>, and *Murusraptor*<sup>25</sup>. Nevertheless, and although this could potentially be due to differences in serial position, the preserved ribs of *Joaquinraptor* appear more gracile than those of *Maip*, and also do not exhibit the prominent, subtriangular medial flange present in the first dorsal rib of that southern Patagonian taxon.

The anterior haemal arch (Fig. 2l) is long and slender, features shared with the enigmatic megaraptoran *Aoniraptor libertatem*<sup>26,62</sup> and *Megaraptor*<sup>3</sup>. The articular surface is saddle-shaped, as in *Aoniraptor*. The haemal canal is ovoid in posterior view, and the blade of the haemal arch curves slightly posteriorly as in *Aoniraptor*, *Megaraptor*, and *Orkoraptor*.

The partially preserved left scapula and coracoid are fused (Fig. 2m), forming a scapulocoracoid. In overall morphology, the partial coracoid resembles the corresponding region of this bone in *Aerosteon*<sup>4,30</sup> and *Maip*<sup>8</sup>, although the posteroventral (=anteroventral, if the long axis of the scapulocoracoid is held horizontally) process of *Joaquinraptor* appears more hooked than in these taxa; moreover, unlike *Maip*, the new form possesses a subglenoid ridge and fossa *sensu*<sup>8,30</sup>. As in *Aerosteon* and *Megaraptor*<sup>3</sup>, the scapular blade of *Joaquinraptor* has sharp dorsal and ventral edges and maintains a near-constant dorsoventral width throughout its length, except at its slightly expanded posterior end. In lateral view, the blade is proportionally wide relative to its length, as in other megaraptorids but contrasting with the



condition in the phylogenetically controversial Patagonian theropod *Gualicho shinyae*, which has a narrower scapular blade<sup>63</sup>.

Although all described megaraptoran humeri are robust, that of *Joaquinraptor* is notably more massive than that of any other member of the group (Fig. 2n, o, p). The shaft is bowed anteriorly as in *Megaraptor*<sup>64</sup>, but in both these genera, the degree of bowing is less marked than in the early diverging megaraptoran *Fukuiraptor* and the Australian megaraptorid *Australovenator*, suggesting that a relatively straight humeral shaft may be a synapomorphy uniting these two Patagonian megaraptorids. Unlike in *Australovenator*, *Fukuiraptor*, and *Megaraptor*, the deltopectoral crest is subrectangular in lateral view (Supplementary Fig. 5). Although, as in these taxa, the crest projects prominently anteriorly and tapers proximally and distally, these features are less marked in *Joaquinraptor*, with the exception of the juvenile *Megaraptor* (MUCPv 595) in which the deltopectoral crest is small (see ref. 6; these authors' Fig. 10).

The left radius (Fig. 2q, r) differs from those of other megaraptorids, particularly in the distal end, which is crescent-shaped (Supplementary Fig. 7), unlike the more ovoid and subtriangular shapes seen in *Australovenator* and *Megaraptor*, respectively. The similarly diagnostic left ulna (Fig. 2s, t) exhibits a well-developed, proximodistally elongate olecranon process that is proportionally longer than that of any other megaraptoran (Supplementary Fig. 6). The bone narrows distally and its shaft is almost straight in lateral and medial views, closely resembling the condition in the *Megaraptor* holotype (MCF-PVPH 79)<sup>1</sup> but not in one of the specimens referred to this taxon (MUCPv 341)<sup>3</sup>; therefore, this feature represents either an autapomorphy of *Joaquinraptor*, a synapomorphy of the former taxon plus *Megaraptor*, or an intraspecifically variable character. There is a relatively marked fossa on the posterolateral surface of the ulnar shaft of *Joaquinraptor*, as in other megaraptorids and, interestingly, some spinosaurids as well<sup>16</sup>. Moreover, there is a notable longitudinal ridge on the medial surface of the proximal end that is not described or illustrated in other megaraptorans, suggesting that it may represent another unique feature of *Joaquinraptor* (Supplementary Fig. 6). The distal end of the ulna is medially expanded and anteroposteriorly compressed, unlike the rounded or triangular shape seen in other megaraptorids<sup>1,3,14,65</sup>.

The right manual ungual II (=manual phalanx II-3; Fig. 2u) resembles those of the Argentinean megaraptorids *Megaraptor* and MCNA-PV 3109<sup>28</sup> (L.M.I. pers. obs.), differing from those of *Australovenator* and early diverging megaraptorans such as *Fukuiraptor* and *Phuwiangvenator*. It is slightly shorter and thicker than in *Australovenator*, a feature that may unite the Argentinean forms to the exclusion of this Australian taxon. Moreover, in *Fukuiraptor*, manual ungual II is much more strongly curved than in megaraptorids (Supplementary Fig. 14).

The nearly complete left femur (Fig. 2v, w) is much more robust than those of *Australovenator* and *Fukuiraptor*, more closely resembling that of the generically unidentified Bajo Barreal Formation megaraptorid UNPSJB-PV 958<sup>7</sup> in this aspect. The posterior flange of the femoral head (*sensu*<sup>5</sup>) present in *Australovenator* and *Fukuiraptor* is lacking in *Joaquinraptor*; moreover, although this may be a taphonomic artifact, the anterior (=lesser) trochanter appears more separated from the remainder of the bone than in these genera. The *Joaquinraptor* femur exhibits an ovoid, projected fourth trochanter that, although slightly more distally placed, closely resembles that of *Australovenator*, differing from the ridge-like structure described in *Fukuiraptor*<sup>2</sup>. The femoral shaft is bowed laterally, unlike the straighter femora of these more ancient megaraptorans.

Much of the right tibia was recovered, albeit surface-collected and broken into several pieces. The shaft is hollow and anteroposteriorly narrow, appearing flat anteriorly but rounded posteriorly. The most striking feature of the distal end (Fig. 2x) is the presence on the anterior surface of a tall, well-marked facet for reception of the ascending process of the astragalus. A tall ascending process of the astragalus

that covers most of the anterodistal surface of the tibia is a diagnostic feature of Coelurosauria<sup>66</sup>.

The distal end of a pedal non-ungual phalanx, probably the right III-3 (Supplementary Fig. 18a–d), has a deep extensor fossa on the dorsal surface, as in *Australovenator*<sup>45</sup>. The medial condyle is slightly taller than the lateral condyle. On both the medial and lateral sides, there are well-marked collateral ligament pits that differ in depth. The right pedal ungual III (=pedal phalanx III-4; Fig. 2y) closely resembles that of *Australovenator*. Another possible partial pedal phalanx, possibly the left IV-1 (Supplementary Fig. 18e–g), is potentially pathological, having an unusually rounded and 'inflated-looking' proximal end in comparison to the same element in *Australovenator*.

**Histological description.** The osteohistological sample of the tibia includes approximately half the complete midshaft section (Supplementary Note 7 and Supplementary Fig. 19a). Despite its incompleteness, the section reveals that the element is formed by a thick cortex surrounding a free medullary cavity. A thick layer of secondarily formed lamellar bone tissue surrounds the medullary cavity (Supplementary Fig. 19b–d). This layer, interpreted as an inner circumferential layer (ICL), contains several radially oriented simple vascular canals, which are open to the medullary cavity (Supplementary Fig. 19b, c). Highly vascularized primary bone tissue predominates in the compacta. The spatial arrangement of intrinsic fibers of the matrix grades from low (i.e., woven-fibered bone) to high (i.e., parallel-fibered bone) (Supplementary Fig. 19e–h). Parallel-fibered bone predominates in the outer portion of the compacta. This tissue is also formed in other areas of the cortex, including the middle and inner regions (Supplementary Fig. 19h). Vascularization is profuse in the entire compacta. Primary osteons are mostly organized in a plexiform pattern (Supplementary Fig. 19d, e, g). This pattern, however, changes in some areas. For example, a distinct layer of disorganized oblique and longitudinal canals is observed in some areas of the middle cortex (Supplementary Fig. 19i). Both the density and the sizes of the lumina of the vascular canals tend to decrease toward the outer edge of the bone. Cyclical growth marks (CGMs) in the form of single and double lines of arrested growth (LAGs) are recorded from the inner to the outer cortex of the sample (Supplementary Fig. 19d, g, j). A minimum number of 19 LAGs can be distinguished. A strong reduction in the spacing between LAGs occurs from the fifth to the 16<sup>th</sup> and from the 17<sup>th</sup> to the 19<sup>th</sup> growth marks (counting from the inner to the outer cortex). Sharpey's fibers are distinct in the outer cortex, in some instances reaching the inner region (Supplementary Fig. 19k). These extrinsic fibers are almost perpendicularly oriented with regard to the subperiosteal surface. Regarding secondary bone, Haversian osteons are mostly formed in the inner cortex (Supplementary Fig. 19b, l). Two particular features are observed with regard to the distribution of secondary osteons: first, concentric patches of these osteons are formed in different levels of the cortex (Supplementary Fig. 19m); and second, despite the fact that these osteons predominate in the inner cortex, they are not formed in the ICL.

The histologically sampled dorsal rib exhibits an anteroposteriorly elongate cross-section. The medial side is mostly flat, but the lateral side is strongly convex (Supplementary Fig. 20a). Whereas one of the margins (it is not possible to determine if this is the anterior or the posterior margin) is broken, the other is preserved and exhibits a groove. The shaft is formed by a distinct cortex of compact bone that surrounds a medullary region occupied by a large pneumatic cavity (Supplementary Fig. 20a). The medial cortex is thicker than the lateral cortex. The compact bone is mostly formed by dense Haversian bone (Supplementary Fig. 20b, c). Secondary osteons occupy almost the entire medial cortex and most of the lateral cortex. The relative sizes and shapes of the secondary osteons are variable. More than one Haversian canal is observed in several secondary osteons. Primary bone tissue is preserved in the outer portion of the lateral cortex and is

formed by well-vascularized fibrolamellar and parallel-fibered bone tissue (Supplementary Fig. 20d–g). Vascular canals are mainly longitudinally and obliquely oriented (Supplementary Fig. 20d–g). Radial canals are also scattered in the compacta. As observed in the tibia, the vascular canal density and the relative size of the canal lumina tend to decrease toward the subperiosteal margin. At least 13 LAGs are recorded in this portion of the cortex (Supplementary Fig. 20d–f). There is no clear pattern regarding the spacing between LAGs. Sharpey's fibers are observed in different areas of the outer cortex, being more abundant in an area located in the lateral portion of the rib (Supplementary Fig. 20h). These extrinsic fibers are also recorded in the thin layer of subperiosteal tissue preserved in the medial side of the rib cortex. The pneumatic cavity is surrounded by a thin layer of secondarily formed lamellar bone tissue, which corresponds to an ICL (Supplementary Fig. 20c, i). Most of this ICL has been remodeled. A distinct patch with Sharpey's fibers is observed in the ICL (Supplementary Fig. 20j, k). These extrinsic fibers are loosely grouped and exhibit two main orientations, which are oblique to the ICL surface.

**Interpretation of ontogenetic stage and minimum age.** Bone tissues preserve records of important events in the life history of a given vertebrate individual<sup>67–69</sup>. In this regard, discrete microstructural changes occur when individuals reach both somatic (i.e., asymptotic growth) and sexual maturity<sup>67,68</sup>. Somatic maturity is usually inferred on the basis of the presence of an external fundamental system (EFS) in the subperiosteal cortex<sup>67,68,70</sup>. An EFS is a form of bone microstructure sometimes observed in the subperiosteal cortex within a transverse bone cross-section, particularly within long bones. This structure is poorly vascularized (or avascular) and is composed of slowly deposited parallel-fibered or lamellar tissue<sup>67,71</sup>. Regarding sexual maturity, it is broadly assumed that this occurred well before somatic maturity in non-avian dinosaurs (e.g.,<sup>72–74</sup>). Sexual maturity is usually inferred on the basis of a distinct change in the spatial organization of intrinsic collagenous fibers (from poorly to well-organized) and/or a noticeable reduction in the spacing between successive CGMs (e.g.,<sup>75,76</sup>). Since there is no evidence for a distinct EFS in the tibia and dorsal rib of the *Joaquinraptor* holotype, we infer that this individual died before the attainment of somatic maturity. It is worth noting, however, that this ontogenetic stage estimate is tentative. In this regard, although post-mortem destruction of the EFS in both skeletal elements seems improbable, this possibility cannot be entirely ruled out. On the other hand, the noticeable reduction in the spacing between LAGs in the tibia suggests that the individual had attained sexual maturity. Assuming that LAGs correspond to annually deposited structures, a minimum age of 19 years is inferred for this *Joaquinraptor* specimen.

The histological analysis of two different skeletal elements of *Joaquinraptor* is important to determine the ontogenetic stage and minimum age of the individual; moreover, the results also provide data regarding which bones are most reliable for minimum age estimation in non-avian dinosaurs. Although the ontogenetic stages of non-avian dinosaurs have commonly been inferred from long bone histology (e.g.,<sup>77–79</sup>), dorsal ribs have recently been demonstrated to be useful for at least minimum age estimation when long bones are not available (i.e.,<sup>80–82</sup>). In megaraptorans, the long bones (in the case of *Joaquinraptor*, the tibia) appear to preserve a better growth record than the dorsal ribs. The lower number of CGMs in the dorsal rib is related to two main factors. First, this element shows a prominent pneumatic cavity in the sampled area. In addition, the dorsal rib of *Joaquinraptor* exhibits a higher degree of secondary remodeling than does the tibia. The presence of a pneumatic cavity and profuse secondary remodeling results in poor preservation of primary bone tissue and, hence, of the growth record in the cortex. A similar condition has been reported in other non-avian dinosaurs, including abelisaurid theropods and titanosaurian sauropods<sup>83–85</sup>.

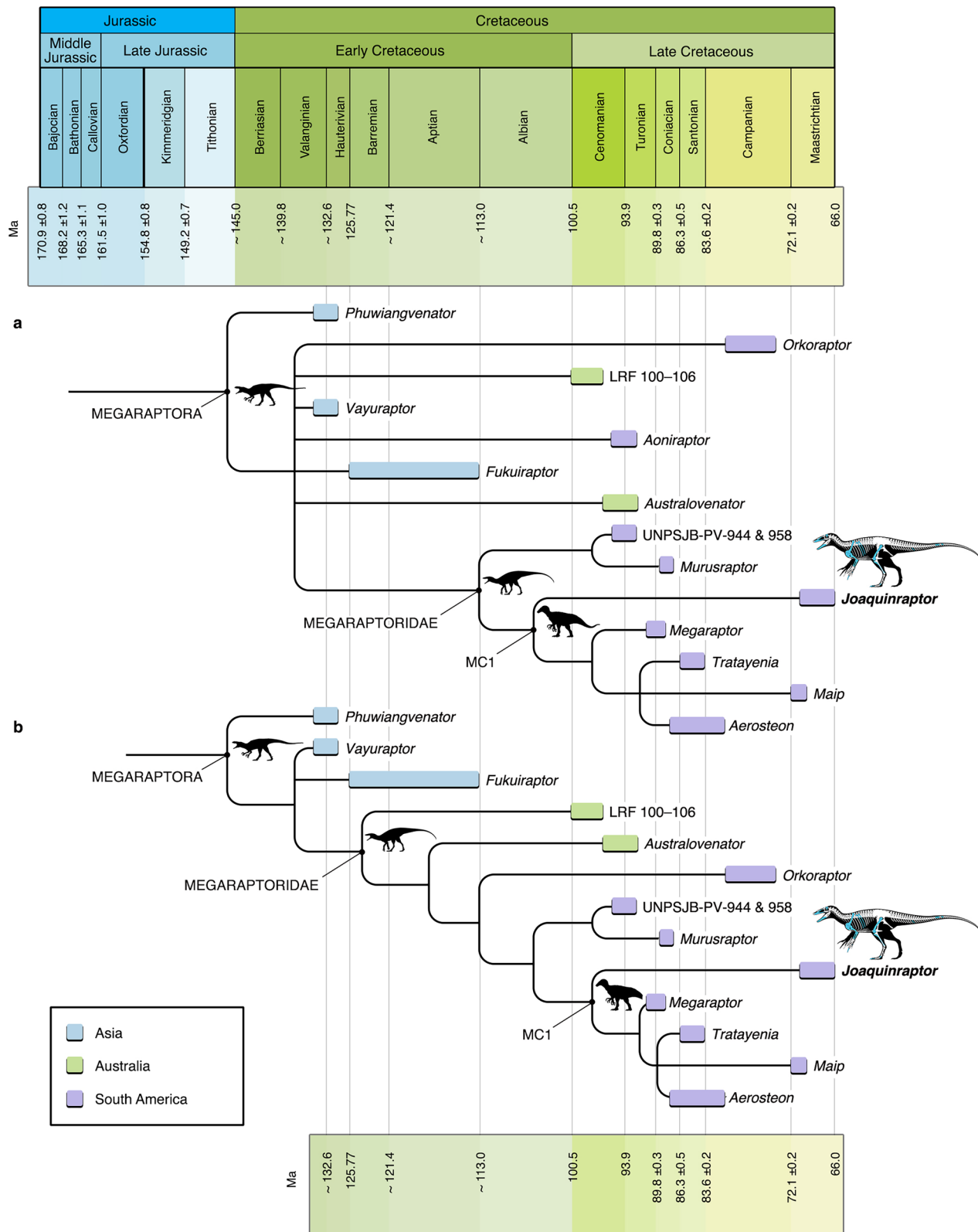
**Histological evidence for postcranial pneumaticity.** Postcranial skeletal pneumaticity (PSP) has been recognized in pterosaurs and diverse clades of saurischian dinosaurs (e.g., refs. 86–89). Among non-avian theropods, PSP is particularly extensive in megaraptorans (e.g., refs. 4,7,8,21,30,62), in which pneumatic cavities reach the mid-caudal vertebrae and selected appendicular elements (e.g., furcula, ilium; see refs. 4,21). Aranciaga Rolando et al.<sup>62</sup> showed that, in *Aoniraptor*, anatomical features related to PSP (i.e., presence of cortical foramina connected to internal cavities) co-occur with histological structures that have also been proposed as evidence for PSP. Specifically, the endosteal bone that forms the internal pneumatic cavities of a caudal centrum and a haemal arch of *Aoniraptor* contains tissue identified as pneumosteon (*sensu*<sup>90</sup>), a bone tissue described in pneumatic structures of some groups of saurischian dinosaurs that is composed of abundant, delicate extrinsic fibers.

The histological analysis conducted on the *Joaquinraptor* dorsal rib provides data regarding the possible correlation of PSP and bone microstructure. As recorded in other saurischians (e.g., refs. 62,90,91), the internal pneumatic cavities are encased by secondarily deposited lamellar bone tissue. Whereas a portion of the secondary lamellar bone tissue does not exhibit any trace of pneumosteon-like tissue (Supplementary Fig. 20i), extrinsic fibers are distinct in other parts of the sample. However, the tissue described here for *Joaquinraptor* (Supplementary Fig. 20j, k) is not identical to the pneumosteon described in other dinosaurs (e.g.,<sup>62,90</sup>). Pneumosteal tissue has been characterized as densely packed, thin, parallel fibers<sup>90</sup>. Conversely, the extrinsic fibers recorded in the dorsal rib of *Joaquinraptor* are loosely packed and exhibit variable orientations, which raises doubts regarding their possible homology with pneumosteon. We conclude that: (1) at least in megaraptorans, PSP is not always correlated with the presence of pneumosteon, as revealed by the absence of this tissue from most of the secondary lamellar bone that surrounds the pneumatic cavity of the rib; and (2) if the extrinsic fibers observed in the endosteal bone of *Joaquinraptor* are homologous to pneumosteal tissue, then this tissue exhibits a higher histological diversity than has previously been proposed.

**Phylogenetic analysis and results.** Two iterative phylogenetic analyses were performed to test the systematic position of *Joaquinraptor*. Because the primary aim of these analyses was to evaluate the systematic position of the new Lago Colhué Huapi Formation taxon within Megaraptoridae—as opposed to resolving the higher-level affinities of Megaraptora among Tetanurae (see, for example<sup>25</sup>)—we added *Joaquinraptor* to the data matrix published by Aranciaga Rolando et al.<sup>8</sup>, which includes the most comprehensive megaraptoran sample of any phylogenetic dataset yet published (Supplementary Notes 8 and 9). Herein, and following Novas et al.<sup>34</sup>, Megaraptora is defined as a stem-based clade containing all taxa closer to *Megaraptor namunhuaiquii* than to *Chilantaisaurus tashuikouensis*, *Neovenator salerii*, *Carcharodontosaurus saharicus*, *Allosaurus fragilis*, *Baryonyx walkeri*, *Tyrannosaurus rex*, and *Passer domesticus*. Megaraptoridae is defined as the most inclusive clade containing *Megaraptor namunhuaiquii* but not *Fukuiraptor kitadaniensis*, following Kotevski et al.<sup>18</sup> and Novas et al.<sup>34</sup>.

Phylogenetic analysis 1 included all operational taxonomic units within the matrix, whereas analysis 2 excluded the fragmentary taxon *Aoniraptor*. Both analyses postulated Megaraptora as well nested within Coelurosauria as the sister group of Tyrannosauroidae (Supplementary Note 10 and Supplementary Fig. 21). Analysis 1 (Fig. 3a) found 102 most parsimonious trees (MPTs) of 1477 steps, with a Consistency Index (CI) of 0.303 and a Retention Index (RI) of 0.594. In the strict consensus of these MPTs, the Thai taxon *Phuwangvenator* is placed at the base of Megaraptora, followed by a polytomy that includes the Asian forms *Vayuraptor* and *Fukuiraptor*, *Australovenator* and the fragmentary specimen LRF 100–106<sup>16</sup> from Australia,





*Orkoraptor* and *Aoniraptor* from Argentine Patagonia, and a clade containing all remaining South American megaraptorans included in the analysis (=Megaraptoridae following the phylogenetic definition employed herein). Relationships among this latter clade are generally well resolved, albeit with low Bremer support values (=1). More specifically, the generically unidentified megaraptorid from the Bajo Barreal Formation (represented by specimens UNPSJB-PV 944 and

UNPSJB-PV 958; see ref. 7) is recovered as the sister taxon of *Murusraptor*. Relative to those of other Patagonian Upper Cretaceous stratigraphic units, the dinosaur fauna of the Bajo Barreal Formation is frequently regarded as being comprised by comparatively plesiomorphic representatives of its constituent clades<sup>31,92</sup>; therefore, the early branching position of the Bajo Barreal taxon within Megaraptoridae is not unexpected. Both these Patagonian forms in turn

**Fig. 3 | Stratigraphically calibrated phylogenetic topologies showing hypothesized relationships of *Joaquinraptor casali* gen. et sp. nov. to other megaraptorans. **a** Strict consensus of most parsimonious trees (MPTs) recovered by analysis 1, which included all taxa in the phylogenetic data matrix. **b** Strict consensus of MPTs recovered by analysis 2, which was conducted after pruning *Aoniraptor libertatem*. Note general geographic and stratigraphic congruence of taxa in (**b**), with earliest diverging megaraptorans known from the Early Cretaceous (Valanginian–Aptian) of Asia (Thailand and Japan), earliest diverging megaraptorids from the mid-Cretaceous (Cenomanian–Turonian) of Australia (New South Wales**

and Queensland), and derived megaraptorids from South America (Argentinean Patagonia), mostly from the Late Cretaceous (Coniacian–Maastrichtian). Abbreviation: MC1 Megaraptorid Clade 1. Silhouettes at selected nodes by T. Tischler (Megaraptora; <https://www.phylopic.org/images/c6099e64-f64e-4685-89a8-10185713ba65/australovenator-wintonensis>), Jagged Fang Designs (Megaraptoridae, <https://www.phylopic.org/images/236a6e22-f960-46c2-acf1-3fc23c38a60a/aerosteon-riocoloradensis>), and T. Dixon (MC1; <https://www.phylopic.org/images/df11d20b-642f-4af2-b524-7a6c073a29b7/maip-macrothorax>), respectively.

constitute the sister taxon of a diverse clade of derived South American megaraptorids. This clade, herein informally termed Megaraptorid Clade 1 (=MC1) for ease of discussion, is united by the ambiguous synapomorphy of the parietals being narrow between the supratemporal fossae, with each reduced to about 10–30% the mediolateral width of the fossa (character 265, state 1; present in *Joaquinraptor* and *Megaraptor*). Within MC1, *Joaquinraptor* is placed as an early diverging form.

Analysis 2 (Fig. 3b), conducted after *Aoniraptor* was pruned, yielded only 12 MPTs of 1476 steps, with a CI of 0.307 and an RI of 0.602. The resulting strict consensus tree shows improved resolution, particularly at the base of Megaraptora. *Phuwiangvenator* is recovered in the same position as in analysis 1, but the abovementioned polytomy is largely resolved, leaving only *Vayuraptor* and *Fukuiraptor* as the next-latest diverging megaraptorans. The Australian forms LRF 100–106 and *Australovenator* are postulated as early diverging megaraptorids and successive sister taxa to a clade comprising all South American megaraptorids in the analysis. This latter group is united by the absence of mesial denticles (character 2, state 2). The recovery of this clade—consisting entirely of Argentinean megaraptorids reaching or exceeding approximately 7 m in total body length—supports the hypothesis that Megaraptoridae experienced a substantial increase in body size upon their arrival in South America during the mid-Cretaceous<sup>7</sup> (see also below). The results of analysis 2 postulate *Orkoraptor* as the earliest diverging member of the Argentinean clade. As in analysis 1, the remaining South American megaraptorids are situated within a clade that is itself divided into two subclades, UNPSJB-PV 944 & 958 + *Murusraptor* and MC1. As in analysis 1, *Joaquinraptor* is recovered as the earliest diverging member of MC1.

In order to down-weight characters that exhibit widespread homoplasy, we applied extended implied weighting ( $k = 3$ ) to both analyses. Similar results were obtained for analysis 1, but (after pruning *Aoniraptor*) analysis 2 yielded better resolution within Megaraptoridae (see Supplementary Fig. 21c). Among derived megaraptorids, both the clades UNPSJB-PV 944 & 958 + *Murusraptor* and MC1 were again recovered; however, there is improved resolution among derived members of MC1, with *Maip* being postulated as the sister taxon of *Tratayenia rosalesi* + *Aerosteon*.

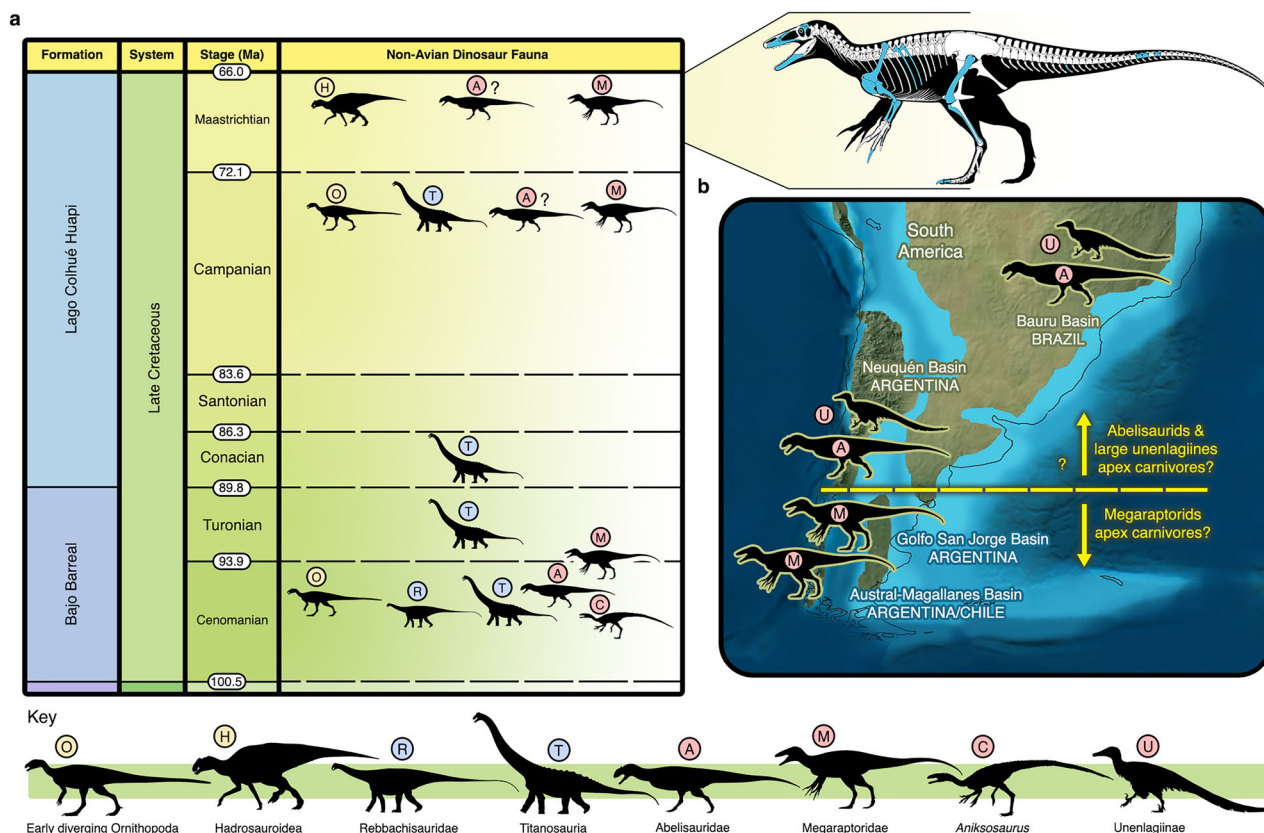
In accordance with the study of Aranciaga Rolando et al.<sup>8</sup>, both our phylogenetic analyses (analysis 1 and analysis 2) recovered nested clades of South American megaraptorids, one more inclusive than the other. Here, we compare the results of our analysis 2 (i.e., our analysis that excluded *Aoniraptor*) to those of the analysis without *Aoniraptor* and the Bajo Barreal Formation megaraptorid (the latter represented by specimens UNPSJB-PV 944 and UNPSJB-PV 958<sup>7</sup>) presented by Aranciaga Rolando et al.<sup>8</sup>. The topologies of the strict consensus trees recovered by these two analyses differ with regard to the hypothesized placements of selected Patagonian megaraptorids. In our analysis, *Megaraptor* occupies a relatively derived position, whereas *Orkoraptor* is supported as an earlier diverging form. Therefore, instead of the clade *Megaraptor* + *Murusraptor* + “Clade B” recovered by Aranciaga Rolando et al.<sup>8</sup> (i.e., these authors’ “Clade A”), we postulate *Orkoraptor* as the outgroup to an unnamed clade that itself includes the sister taxa UNPSJB-PV 944 & 958 + *Murusraptor* and MC1 (the latter of which encompasses all megaraptorid taxa within Aranciaga Rolando et al.’s<sup>8</sup>

“Clade B,” with the addition of *Joaquinraptor* and the aforementioned exception of *Orkoraptor*).

In sum, our phylogenetic analyses yielded several notable results that may be pertinent to future discussions of megaraptoran systematics. First, and unsurprisingly given the data matrix we used, the analyses support the placement of Megaraptora within Coelurosauria, as the sister group of Tyrannosauroidae. Moreover, *Joaquinraptor* is recovered as the earliest diverging member of a newly identified clade of derived South American megaraptorids, herein termed Megaraptorid Clade 1 (=MC1). Nevertheless, although the position of *Joaquinraptor* within Megaraptoridae on the whole is well supported, MC1 is presently supported by only a single synapomorphy; furthermore, two putative members of the clade (*Maip* and *Tratayenia*) are relatively incompletely preserved, being presently known only from postcranial (mainly postcranial axial skeletal) bones. As such, future analyses should evaluate the proposed monophyly and taxonomic content of MC1 and, therefore, its potential systematic validity, as well as the specific relationships of taxa within the clade.

**Implications for Patagonian theropod faunal provincialism and megaraptorid palaeobiology.** The post-Turonian central (Fig. 4a) and southern Patagonian fossil record of large-bodied non-avian theropod dinosaurs is, at the moment, numerically dominated by megaraptorids<sup>7,8,22,23,31,53</sup>. Nevertheless, to date, all described central and southern Patagonian megaraptorid specimens have been fragmentary; therefore, *Joaquinraptor* is important in being, by a substantial margin, the most complete representative of the clade yet found from this geographic area. Moreover, this taxon was recovered from only a few meters below the top of the Lago Colhué Huapi Formation, and is therefore Maastrichtian (probably late Maastrichtian) in age, rendering it the latest surviving megaraptoran yet known from central Patagonia and one of the stratigraphically youngest (quite possibly the youngest) members of the group found anywhere in the world. Although more evidence is needed, Patagonian megaraptorids present a suite of putative synapomorphies, such as the absence of mesial denticles from the tooth crowns, a manual ungual I that is subequal in length to the ulna, and a well-developed “mediolateral platform” on the flexor tubercle of this ungual<sup>8,35</sup>, that suggest that these theropods may have been evolutionarily and geographically differentiated from earlier diverging members of the group (e.g., *Australovenator*).

Furthermore, the discovery of *Joaquinraptor* supports the hypothesis that megaraptorids were the apex predators and the most abundant large-bodied theropods in the latest Cretaceous (Campanian–Maastrichtian) of central and southern Patagonia, contrasting with penecontemporaneous faunas from more northerly areas of South America that appear to have been dominated by abelisaurids and large unenlagiines (Fig. 4b; see also refs. 7,8,23,93). Novas et al.<sup>23</sup> suggested that this putative theropod provincialism may have been related to palaeoenvironmental differences. Notably, however, the sedimentology of the middle (i.e., Santonian–?lower Maastrichtian) section of the Lago Colhué Huapi Formation, which has yielded several other megaraptorid fossils<sup>31,55</sup>; (Fig. 4a), is characterized by fluvial channel morphologies that indicate that the fauna it preserves lived inland, well away from the palaeoshoreline; moreover, clay analyses support a seasonally dry or semiarid climate for this part of the



**Fig. 4 | Dinosaur fauna of the Chubut Group and hypothesized distribution of theropod apex predators in the latest Cretaceous of southern South America.**

**a** Upper Cretaceous non-avian dinosaur faunas of the Chubut Group, Golfo San Jorge Basin, central Patagonia, Argentina, indicating stratigraphic position of *Joaquinraptor casali* gen. et sp. nov. (denoted by skeletal reconstruction).

**b** Hypothesized distribution of medium- to large-bodied theropod dinosaurs occupying apex predator niches in selected South American depositional basins during the Campanian and Maastrichtian stages of the Late Cretaceous. Note absence of Megaraptoridae north of the Golfo San Jorge Basin. Palaeogeographic basemap ©2023 Colorado Plateau Geosystems Inc., used with permission. Dinosaurian silhouettes except Megaraptoridae by S. Hartman (early diverging

Ornithopoda, <https://www.phylopic.org/images/05b2229f-34dd-4fdc-ae6b-8158a6b38119/parksosaurus-warreni>; Hadrosauroidae, <https://www.phylopic.org/images/4ae90b8f-bbb4-47cd-a914-98e1c845cfe/gryposaurus-monumentensis>; Rebbachisauridae, <https://www.phylopic.org/images/a5b8df09-f5f9-43ae-ae2a-7822ff3e3d56/nigersaurus-taqueti>; Titanosauria, <https://www.phylopic.org/images/33ae8000-0046-41bb-b68a-189a0eed3eca/opisthocoeleicaudia-skarzynskii>; Abelisauridae, <https://www.phylopic.org/images/d82ad39a-0770-4f85-99a6-fd17bd1bf64/majungasaurus-crenatissimus>), FunkMonk (*Aniksosaurus*, <https://www.phylopic.org/images/d3ae8c0d-107d-45de-992e-177c551e079d/coelurus-agilis>), and T. Dixon (Unenlagiinae, <https://www.phylopic.org/images/eb481f28-4f1a-4cfc-96ea-f3ab0a622e6b/austroaptor-cabazai>).

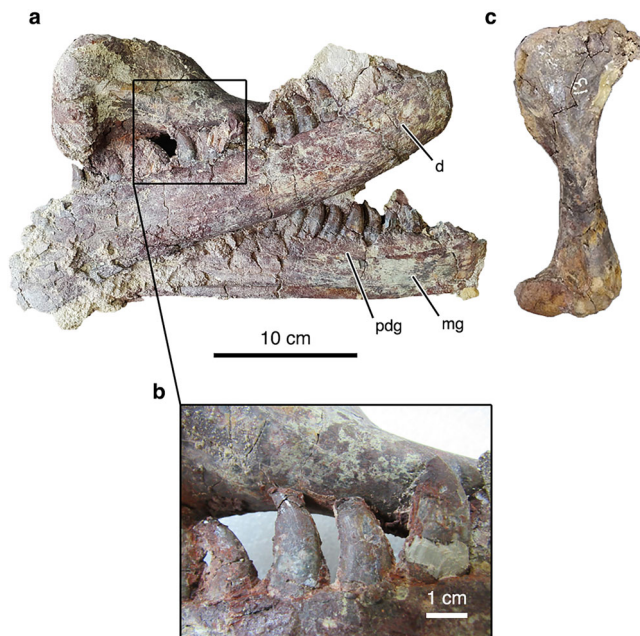
formation (see ref. 46 and references therein). Conversely, in the uppermost section of the formation, at a stratigraphic level close to that from which *Joaquinraptor* was collected, sedimentological and palynological inferences support the existence of a warm, humid floodplain palaeoenvironment close to the sea (see Supplementary Note 3). Consequently, it appears that, at least in central Patagonia, megaraptorids endured significant palaeoclimatic changes and successfully adapted to fluctuating palaeoenvironmental conditions through Late Cretaceous time, raising the possibility that these theropods were able to thrive in a broad range of habitats. As such, alongside putative palaeoenvironmental distinctions, additional factors, such as the potential palaeogeographic isolation of central and southern Patagonia from more northern areas of South America by epicontinental seas (Fig. 4b; see also, e.g., Reguero and Goin<sup>94</sup> [these authors' Fig. 8]) and/or pronounced floristic differences (see, for example, the reconstructed Late Cretaceous palynological bioprovinces proposed by Vajda and Bercovic<sup>95</sup> [these authors' Fig. 3b]) may also have driven the differentiation of large-bodied theropod faunas in southern South America at the end of the Mesozoic.

The distinctive megaraptorid forelimb may have played an important role in the apparent adaptability of these theropods, perhaps allowing them to exploit resources in varying ways and/or to feed

on an array of vertebrate taxa. For example, megaraptorids may have used their hypertrophied manus with its enlarged, trenchant unguals to access soft tissues<sup>96</sup> and/or to aid in prey capture and manipulation<sup>9,20,25,97,98</sup>. In addition, several features of the megaraptorid antebrachium, such as the proximodistally elongate olecranon process (the insertion area of *M. triceps brachii*) and the vertical ridge on the proximolateral surface of the ulna (a probable osteological correlate for the *M. anconeus* insertion<sup>9,99,100</sup>), may have conferred enhanced extension and flexion capabilities.

As mentioned above, in addition to the *Joaquinraptor* skeleton, a crocodyliform right humerus was recovered from the type quarry of the new megaraptorid. Interestingly, this bone was situated between the closely associated left and right dentaries of the *Joaquinraptor* holotype, with the apices of several tooth crowns of the megaraptorid in direct contact with the humeral shaft, which also shows potential tooth marks (Fig. 5); moreover, this humerus is the only non-megaraptorid vertebrate fossil yet identified from the site. As such, this discovery may constitute direct evidence regarding prey selection within Megaraptoridae (although isolated megaraptorid teeth have been found in association with other saurischian dinosaur specimens, potentially also indicating feeding behavior [e.g., refs. 41,101]). Nevertheless, this megaraptorid/crocodyliform co-occurrence could





**Fig. 5 | Putative evidence of dietary preference and/or prey capture in *Joaquinraptor casali* gen. et sp. nov.** **a** Dentaries of *Joaquinraptor* (UNPSJB-PV 1112) and an associated crocodyliform right humerus (UNPSJB-PV 1113) in the position in which they were recovered prior to their mechanical separation in the laboratory (right dentary in lateral view, left dentary in medial view, humerus in posterior view [proximal end only]). **b** Detail of **a** showing a portion of the crocodyliform humeral shaft (UNPSJB-PV 1113) in contact with the right dentary tooth crowns of *Joaquinraptor* (UNPSJB-PV 1112). **c** Crocodyliform right humerus (UNPSJB-PV 1113) in anterior view. Abbreviations: d dentary, mg Meckelian groove, pdg paracanth groove.

conceivably be instead related to other behaviors, such as interspecific combat or competition for food, or even just a coincidental association (e.g., the humerus was moved to a position between the *Joaquinraptor* dentaries by water currents). Detailed taphonomic studies are currently underway to assess the probable cause and potential significance of this association.

In sum, the singular morphology of the megaraptorid forelimb may have played an important role in the evolutionary success of these theropods in the Late Cretaceous of central and southern Patagonia. The specialized development and capabilities of the forelimb were almost certainly adaptations related to prey capture<sup>9,97,98</sup>, and, in the case of *Joaquinraptor*, these prey items may have at least occasionally included crocodyliforms (Fig. 5).

**Megaraptorid palaeobiogeography and body size evolution.** Based on the fossil record of Megaraptoridae as it is currently understood, Lamanna et al.<sup>7</sup> hypothesized that South American members of the clade underwent a substantial increase in body size with respect to their earlier diverging and mostly stratigraphically older Australian relatives. Specifically, megaraptorids are proposed to have experienced an increase in size from approximately 5 m in total body length and 300 kg in body mass to more than 7 m in length and 1000 kg in mass not long after their dispersal to South America from Australia, presumably via Antarctica, during the mid-Cretaceous<sup>7</sup>. Moreover, South American megaraptorids are hypothesized to have maintained this elevated body size until their extinction at or near the end of the Cretaceous<sup>7,8</sup>.

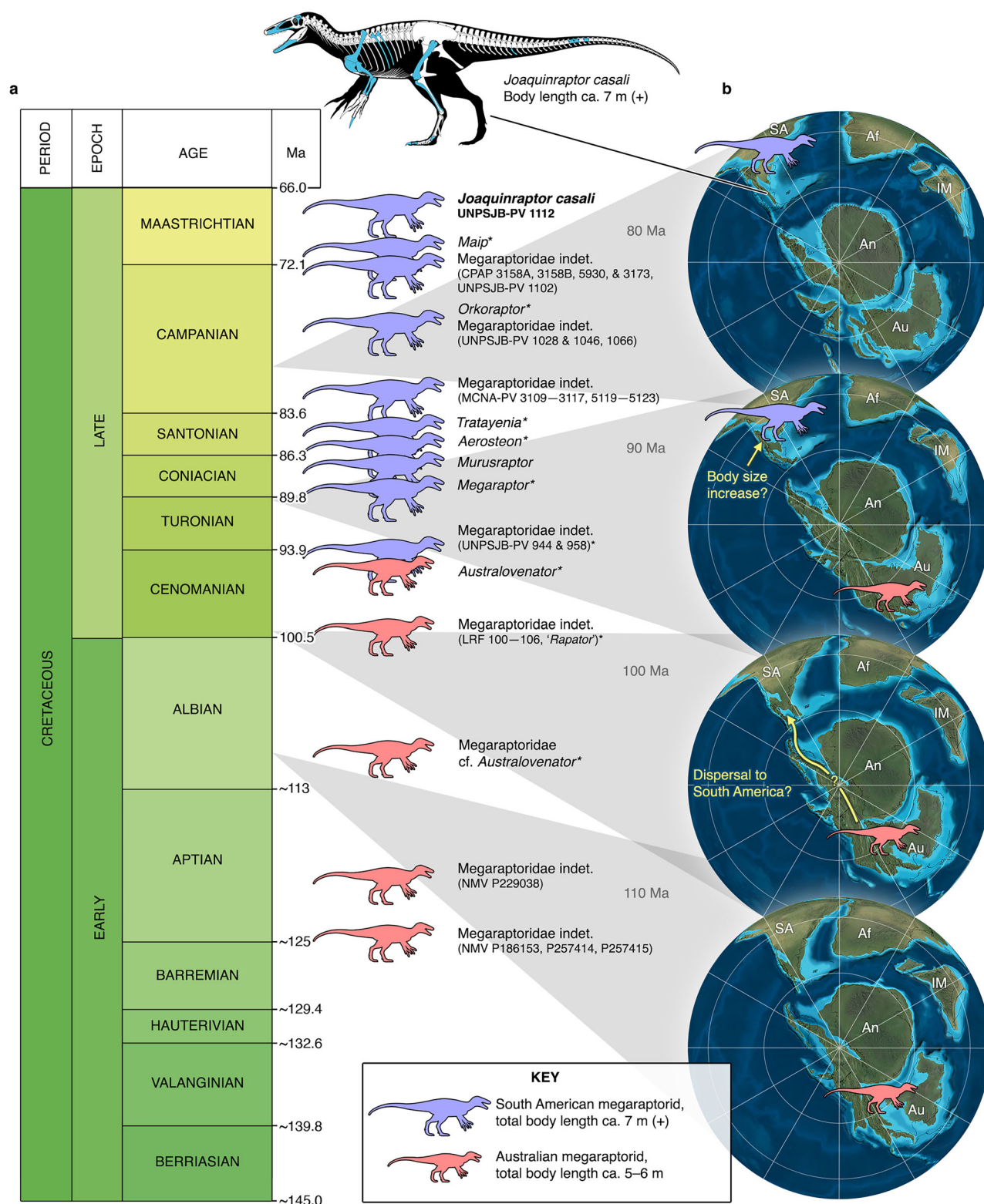
The nearly complete (missing only the medial condyle of the distal end) left femur of the holotype of the Maastrichtian megaraptorid *Joaquinraptor* has a total proximodistal length of 685 mm and a circumference (measured immediately distal to the fourth trochanter) of

275 mm (Table 2). Based on the scaling equation presented by Campione and Evans<sup>102</sup> and employed in many subsequent works (e.g., refs. 7,103), this circumference yields a mass estimate of 1036 kg for this individual. Approximately the same circumference (280 mm) was obtained from the fragmentary femur of another large-bodied megaraptorid specimen from a nearby exposure of the stratigraphically older (Cenomanian–Turonian) Bajo Barreal Formation (UNPSJB-PV 958), from which an estimated body mass of 1089 kg was derived<sup>7</sup>. Furthermore, a mass estimate that is precisely the same as that generated for *Joaquinraptor* (1036 kg) was proposed for MUCPV 1353, an undescribed partial megaraptorid skeleton (probably referable to *Megaraptor*) from the Turonian–Coniacian Portezuelo Formation of northwestern Patagonia<sup>7</sup>. On the other hand, the type specimen of the early diverging Australian megaraptorid *Australovenator* has a femur length of 578 mm<sup>15</sup>, whereas that of the holotype of the non-megaraptorid megaraptoran *Fukuiraptor* is 507 mm<sup>2</sup>. Likewise, the estimated body mass of the former taxon (314 kg<sup>103</sup>) is approximately 3.3–3.5 times less than that of *Joaquinraptor*, MUCPV 1353, and UNPSJB-PV 958<sup>7</sup>. Aranciaga Rolando et al.<sup>8</sup> proposed a body length of more than 9 m for *Maip*, a megaraptorid from the Maastrichtian Chorrillo Formation of southern Patagonia, which would likely establish this taxon as the largest-bodied megaraptorid yet known.

In sum, *Joaquinraptor* accords with Lamanna et al.'s<sup>7</sup> hypothesis of body size evolution in South American megaraptorids in that it was probably longer than 7 m and heavier than 1000 kg; in other words, approximately the same size as MUCPV 1353 and UNPSJB-PV 958. Furthermore, *Joaquinraptor* is from the Maastrichtian—probably the late Maastrichtian—supporting the hypothesis that Megaraptoridae maintained this elevated body size until their extinction (Fig. 6).

Recently, Kotevski et al.<sup>19</sup> proposed that selected megaraptorid postcranial elements (a set of two articulated caudal vertebrae and their corresponding haemal arches [NMV P257414], an isolated partial manual ungual II [NMV P186153], and the distal end of a tibia [NMV P257415]) from the upper Barremian–lower Aptian upper Strzelecki Group of Victoria, Australia suggest that ca. 6 m individuals of this theropod clade inhabited that continent during the Early Cretaceous. The upper Strzelecki Group bone that implies the largest individual—a megaraptorid perhaps slightly greater than 6 m in body length—is the fragmentary manual ungual II NMV P186153, which was also described and/or analysed at length by Benson et al.<sup>104</sup>, Poropat et al.<sup>17</sup>, and White and Campione<sup>105</sup>. That said, this claw is isolated and incomplete, and as such, the hypothesized body size of the individual to which it pertained—though it may well be correct—cannot be considered to be robustly documented given the fragmentary nature of the currently available evidence. Consequently, we consider the hypothesis that >6 m megaraptorids were present in Australia to be only tentatively supported at present. Moreover, even if NMV P186153 does indeed represent a megaraptorid marginally greater than 6 m in body length, the likelihood remains that all definitive South American megaraptorid records indicate animals that are still larger (ca. 7 m [±] in length and ca. 1000 kg [±] in mass). As such, in our view, Lamanna et al.'s<sup>7</sup> hypothesis that megaraptorids experienced a substantial increase in body size shortly after their arrival in South America remains consistent with the presently available data (Fig. 6). That said, it may potentially be only a matter of time before definitive evidence of large-bodied (≥7 m, ≥1000 kg) megaraptorids is discovered in the Early or mid-Cretaceous of Australia.

To conclude, as proposed by Lamanna et al.<sup>7</sup> and supported by more recent discoveries (e.g.,<sup>8,106</sup>), including *Joaquinraptor* (Fig. 6), South American megaraptorids appear to have attained larger body sizes (~7 m or greater in total body length) than their generally more ancient relatives from Australia (~5–6 m<sup>19</sup>). Patagonian megaraptorids may have continued to increase in size through the remainder of their evolutionary history—perhaps spurred at least in part by the extinction of carcharodontosaurids during the Turonian<sup>8,106</sup>—with some latest



**Fig. 6 | Synopsis of fossil record, body size evolution, and palaeobiogeography of Megaraptoridae (modified and updated from Lamanna et al.<sup>7</sup> [these authors' Fig. 13]).** **a** Stratigraphic distribution of definitive megaraptorid fossils through the Cretaceous. Asterisks adjacent to some records indicate that additional, generically indeterminate megaraptorid materials have been reported from the same stratigraphic unit. Stratigraphic position of *Aerosteon riocoloradensis* within the upper Coniacian–lower Campanian is uncertain (see ref. 21). **b** Hypothesis of palaeobiogeography and body size evolution in Megaraptoridae. Megaraptorids are

considered to have originated in eastern Gondwana (Australia) during the Early Cretaceous, then dispersed to western Gondwana (South America) via Antarctica during the mid- (i.e., late Early or early Late) Cretaceous (see refs. 7,8,16). Red silhouette represents Australian megaraptorid ca. 5–6 m in total body length; blue silhouette represents ca. 7 m (+) South American megaraptorid. Cretaceous time-scale in **a** from ref. 110; palaeogeographic reconstructions in **b** are south polar views at (bottom to top) 110, 100, 90, and 80 Ma (©2023 Colorado Plateau Geosystems Inc., used with permission).



Cretaceous forms (e.g., *Maip*) potentially reaching body lengths of 9 m or more<sup>8</sup>. Among other benefits, this may have allowed megaraptorids to occupy the apex predator niches in their respective habitats. Moreover, and as evidenced by the Maastrichtian (probably late Maastrichtian) stratigraphic occurrence of *Joaquinraptor*, these theropods may well have survived to the Cretaceous/Palaeogene boundary, only to ultimately perish alongside all other non-avian dinosaurs.

## Methods

### Field permits and permissions

The field research that recovered the holotypic specimen of *Joaquinraptor casali* (UNPSJB-PV 1112) was carried out at the informally named Valle Joaquín site in south-central Chubut Province, central Patagonia, Argentina. Personnel from the Laboratorio de Paleontología de Vertebrados Dr. Rubén Martínez of the Universidad Nacional de la Patagonia San Juan Bosco obtained all necessary permissions from the Subsecretaría de Cultura de la Provincia del Chubut and the relevant landowner to conduct this fieldwork.

### Palaeohistology

We conducted a histological analysis to assess the ontogenetic stage and other osteohistological attributes of UNPSJB-PV 1112. Transverse sections were obtained from the midshaft of the right tibia and the proximal shaft of a dorsal rib. Histological thin sections were prepared at the petrographic laboratory of the Universidad Nacional de San Luis (San Luis, Argentina). The thin sections were prepared using standard petrographic methods (e.g.,<sup>107</sup>) and studied using a petrographic polarizing microscope (Leica DM750P). The nomenclature and definitions of structures used in this study are derived from Francillon-Vieillot et al.<sup>108</sup> and de Buffrénil and Quilhac<sup>69</sup>.

### Phylogenetic analysis

Morphological characters 1–351 used in our phylogenetic analysis were sourced from Aranciaga Rolando et al.<sup>36</sup> and references therein, with their descriptions as edited and presented by Lamanna et al.<sup>7</sup> (these authors' Supplementary Appendix A). In a few cases, these character descriptions were slightly further modified (without changing their meaning) from those published by Lamanna et al.<sup>7</sup>. Characters 352–357 were subtly edited (also without changing their meaning) from Aranciaga Rolando et al.<sup>8</sup>.

We used the software TNT (Tree analysis using New Technology) version 1.5<sup>109</sup> to evaluate the phylogenetic position of *Joaquinraptor* as well as to test the topology within Megaraptora after the addition of the new theropod. We employed the traditional search function using the tree bisection-reconnection (TBR) swapping algorithm and a random seed of 1. The traditional search included 10,000 replications with ten trees saved per replication. We performed two analyses (analysis 1 and analysis 2) under equally weighted parsimony. Analysis 1 included all megaraptoran taxa analysed by Aranciaga Rolando et al.<sup>8</sup>, but *Aoniraptor libertatem* was pruned from analysis 2. *Aoniraptor* was excluded from analysis 2 because this theropod was the least stable megaraptoran taxon in analysis 1, in part due to its fragmentary preservation (i.e., it is presently known from less than 15% of the skeleton). Bremer support values within Megaraptora were low (=1) in the strict consensus trees generated by both analyses. Subsequently, we applied extended implied weighting ( $k=3$ ) to both analyses to down-weight particularly homoplastic characters.

### Reporting summary

Further information on research design is available in the Nature Portfolio Reporting Summary linked to this article.

### Data availability

The authors declare that all data generated or analysed during this study, which support the results published herein, are included in this

article and its Supplementary Information file. The fossils are housed in the Repositorio de Materiales Científicos y Didácticos Dr. Eduardo Musacchio of the Universidad Nacional de la Patagonia San Juan Bosco (UNPSJB) in Comodoro Rivadavia, Chubut Province, Argentina. Correspondence and requests for material should be addressed to L.M.I. Qualified researchers may access the holotypic specimen of *Joaquinraptor casali* (UNPSJB-PV 1112) and the associated crocodyliform humerus (UNPSJB-PV 1113) by contacting Gabriel Casal of the UNPSJB (paleogac@yahoo.com.ar).

## References

- Novas, F. E. *Megaraptor namunhauquii*, gen. et sp. nov., a large-clawed, Late Cretaceous theropod from Patagonia. *J. Vertebrate Paleontol.* **18**, 4–9 (1998).
- Azuma, Y. & Currie, P. J. A new carnosaur (Dinosauria: Theropoda) from the Lower Cretaceous of Japan. *Can. J. Earth Sci.* **37**, 1735–1753 (2000).
- Calvo, J. O., Porfiri, J. D., Veralli, C., Novas, F. & Poblete, F. Phylogenetic status of *Megaraptor namunhauquii* Novas based on a new specimen from Neuquén, Patagonia, Argentina. *Ameghiniana* **41**, 565–575 (2004).
- Sereno, P. C. et al. Evidence for avian intrathoracic air sacs in a new predatory dinosaur from Argentina. *PLoS ONE* **3**, e3303 (2008).
- Hocknull, S. A. et al. New mid-Cretaceous (latest Albian) dinosaurs from Winton, Queensland, Australia. *PLoS ONE* **4**, e6190 (2009).
- Porfiri, J. D. et al. Juvenile specimen of *Megaraptor* (Dinosauria, Theropoda) sheds light about tyrannosauroid radiation. *Cretac. Res.* **51**, 35–55 (2014).
- Lamanna, M. C., Casal, G. A., Martínez, R. D. F. & Ibiricu, L. M. Megaraptorid (Theropoda: Tetanurae) partial skeletons from the Upper Cretaceous Bajo Barreal Formation of central Patagonia, Argentina: implications for the evolution of large body size in Gondwanan megaraptorans. *Ann. Carnegie Mus.* **86**, 255–294 (2020).
- Aranciaga Rolando, A. M. et al. A large Megaraptoridae (Theropoda: Coelurosauria) from Upper Cretaceous (Maastrichtian) of Patagonia, Argentina. *Sci. Rep.* **12**, 6318 (2022).
- Aranciaga Rolando, A. M. et al. Reconstruction of the pectoral girdle and forelimb musculature of Megaraptora (Dinosauria: Theropoda). *Anat. Rec.* **306**, 1804–1823 (2023).
- Currie, P. J. & Azuma, Y. New specimens, including a growth series, of *Fukuiraptor* (Dinosauria, Theropoda) from the Lower Cretaceous Kitadani Quarry of Japan. *J. Paleontol. Soc. Korea* **22**, 173–193 (2006).
- Samathi, A., Chanthasit, P. & Sander, P. M. Two new basal coelurosaurian theropod dinosaurs from the Lower Cretaceous Sao Khua Formation of Thailand. *Acta Palaeontol. Polonica* **64**, 239–260 (2019).
- Samathi, A., Suteethorn, S., Pradit, N. & Suteethorn, V. New material of *Phuwiangvenator yaemniyomi* (Dinosauria: Theropoda) from the type locality: implications for the early evolution of Megaraptora. *Cretac. Res.* **131**, 105093 (2022).
- Samathi, A., Weluwanarak, J., Duanyai, P., Kaikaew, S. & Suteethorn, S. An unusual metatarsal of theropod dinosaur from the Lower Cretaceous of Thailand: the first detailed study of paleopathology in Megaraptora. *Historical Biol.* **36**, 467–472 (2024).
- Smith, N. D. et al. A *Megaraptor*-like theropod (Dinosauria: Tetanurae) in Australia: support for faunal exchange across eastern and western Gondwana in the mid-Cretaceous. *Proc. R. Soc. Lond. B* **275**, 2085–2093 (2008).
- White, M. A. et al. New *Australovenator* hind limb elements pertaining to the holotype reveal the most complete neovenatorid leg. *PLoS ONE* **8**, e68649 (2013).
- Bell, P. R., Cau, A., Fanti, F. & Smith, E. T. A large-clawed theropod (Dinosauria: Tetanurae) from the Lower Cretaceous of Australia



- and the Gondwanan origin of megaraptorid theropods. *Gondwana Res.* **36**, 473–487 (2016).
17. Poropat, S. F., White, M. A., Vickers-Rich, P. & Rich, T. H. New megaraptorid (Dinosauria: Theropoda) remains from the Lower Cretaceous Eumeralla Formation of Cape Otway, Victoria, Australia. *J. Vertebrate Paleontol.* **39**, e1666273 (2019).
  18. Kotevski, J. et al. A megaraptorid (Dinosauria: Theropoda) frontal from the upper Strzelecki Group (Lower Cretaceous) of Victoria, Australia. *Cretac. Res.* **154**, 105769 (2024).
  19. Kotevski, J. et al. Evolutionary and paleobiogeographic implications of new carcharodontosaurian, megaraptorid, and unenlagiine theropod remains from the upper Lower Cretaceous of Victoria, southeast Australia. *J. Vertebrate Paleontol.* **44**, e2441903 (2025).
  20. Porfiri, J. D., dos Santos, D. & Calvo, J. O. New information on *Megaraptor namunhuaiquii* (Theropoda: Tetanurae), Patagonia: considerations on paleoecological aspects. *Arq. Mus. Nac.* **65**, 545–550 (2007).
  21. Porfiri, J. D., Juárez Valieri, R. D., dos Santos, D. D. & Lamanna, M. C. A new megaraptoran theropod dinosaur from the Upper Cretaceous Bajo de la Carpa Formation of northwestern Patagonia. *Cretac. Res.* **89**, 302–319 (2018).
  22. Novas, F. E., Ezcurra, M. D. & Lecuona, A. *Orkoraptor burkei* nov. gen. et sp., a large theropod from the Maastrichtian Pari Aike Formation, southern Patagonia, Argentina. *Cretac. Res.* **29**, 468–480 (2008).
  23. Novas, F. E. et al. Paleontological discoveries in the Chorrillo Formation (upper Campanian-lower Maastrichtian, Upper Cretaceous), Santa Cruz Province, Patagonia, Argentina. *Rev. Mus. Argent. Cienc. Nat.* **21**, 217–293 (2019).
  24. Paulina-Carabajal, A. & Coria, R. A. An unusual theropod frontal from the Upper Cretaceous of north Patagonia. *Alcheringa Australas. J. Paleontol.* **39**, 514–518 (2015).
  25. Coria, R. A. & Currie, P. J. A new megaraptoran dinosaur (Dinosauria, Theropoda, Megaraptoridae) from the Late Cretaceous of Patagonia. *PLoS ONE* **11**, e0157973 (2016).
  26. Motta, M. J. et al. New theropod fauna from the Upper Cretaceous (Huincul Formation) of northwestern Patagonia, Argentina. *N. Mex. Mus. Nat. Hist. Sci. Bull.* **71**, 231–253 (2016).
  27. Paulina-Carabajal, A. & Currie, P. J. The braincase of the theropod dinosaur *Murusraptor*: osteology, neuroanatomy and comments on the paleobiological implications of certain endocranial features. *Ameghiniana* **54**, 617–640 (2017).
  28. Méndez, A. H., Gianechini, F. A., Canale, J. I. & Díaz-Martínez, I. A new megaraptorid specimen (Theropoda, Coelurosauria) from Cañadón Amarillo (Anacleto Formation, Campanian, Upper Cretaceous), Mendoza Province, Argentina. *Jorn. Argentinas Paleontol. Vertebr.* **33**, R25–R26 (2019).
  29. Méndez, A. H. et al. Megaraptoran remains from Cerro Overo-la Invernada fossil site (Bajo de la Carpa Formation, Santonian), northern Patagonia, Argentina. *Jorn. Paleontol. de la Cuenca Neuquina, Libro de Resúmenes* **3**, R261 (2022).
  30. Aranciaga Rolando, A. M., Méndez, A., Canale, J. & Novas, F. Osteology of *Aerosteon riocoloradensis* (Serenó et al. 2008) a large megaraptoran (Dinosauria: Theropoda) from the Upper Cretaceous of Argentina. *Historical Biol.* **34**, 226–282 (2022).
  31. Ibiricu, L. M., Casal, G. A., Martínez, R. D., Alvarez, B. N. & Poropat, S. F. New materials and an overview of Cretaceous vertebrates from the Chubut Group of the Golfo San Jorge Basin, central Patagonia, Argentina. *J. South Am. Earth Sci.* **98**, 102460 (2020).
  32. Meso, J. G. et al. Shed teeth from Portezuelo Formation at Sierra del Portezuelo reveal a higher diversity of predator theropods during Turonian–Coniacian times in northern Patagonia. *BMC Ecol. Evol.* **24**, 59 (2024).
  33. Benson, R. B. J., Carrano, M. T. & Brusatte, S. L. A new clade of archaic large-bodied predatory dinosaurs (Theropoda: Allosaurioidea) that survived to the latest Mesozoic. *Naturwissenschaften* **97**, 71–78 (2010).
  34. Novas, F. E., Agnolín, F. L., Ezcurra, M. D., Porfiri, J. & Canale, J. I. Evolution of the carnivorous dinosaurs during the Cretaceous: the evidence from Patagonia. *Cretac. Res.* **45**, 174–215 (2013).
  35. Novas, F. E., Aranciaga Rolando, A. M. & Agnolín, F. L. Phylogenetic relationships of the Cretaceous Gondwanan theropods *Megaraptor* and *Australovenator*: the evidence afforded by their manual anatomy. *Mem. Mus. Vic.* **74**, 49–61 (2016).
  36. Aranciaga Rolando, A. M., Novas, F. E. & Agnolín, F. L. A reanalysis of *Murusraptor barrosaensis* Coria & Currie (2016) affords new evidence about the phylogenetical relationships of Megaraptora. *Cretac. Res.* **99**, 104–127 (2019).
  37. Seeley, H. G. On the classification of the fossil animals commonly named Dinosauria. *Proc. R. Soc. Lond.* **43**, 165–171 (1887).
  38. Marsh, O. C. Principal characters of American Jurassic dinosaurs, part V. *Am. J. Sci., Ser. 3* **21**, 417–423 (1881).
  39. Gauthier, J. Saurischian monophyly and the origin of birds. *Mem. Calif. Acad. Sci.* **8**, 1–55 (1986).
  40. Meso, J. G., Baiano, M. A., Canale, J. I. & Salgado, L. Descripción y caracterización de la dentición de los dinosaurios terópodos megaraptores. *Libro de Resúmenes de la Reunión de Comunicaciones de la Asociación Paleontológica Argentina*, 53 (2017).
  41. Meso, J. G. et al. Theropods from the La Bonita site, Bajo de la Carpa Formation (Neuquén Group, Santonian), Río Negro, Argentina: analysis of dental evidence. *Cretac. Res.* **137**, 105250 (2022).
  42. Ghiglione, M. C. et al. Santonian–Campanian continentalization in the Austral-Magallanes Basin: regional correlation, provenance and geodynamic setting. *Cretac. Res.* **128**, 104968 (2021).
  43. Moyano-Paz, D. et al. The uppermost Cretaceous continental deposits at the southern end of Patagonia, the Chorrillo Formation case study (Austral-Magallanes Basin): sedimentology, fossil content and regional implications. *Cretac. Res.* **130**, 105059 (2022).
  44. Novas, F. E. et al. Paleontological discoveries in SW Argentina: new insights into the end of the dinosaur era in South America. *J. Vertebrate Paleontol. Program Abstr.* **2024**, 413 (2024).
  45. Vallati, P., De Sosa Tomas, A. & Casal, G. A Maastrichtian terrestrial paleoenvironment close to the K/Pg boundary in the Golfo San Jorge Basin, Patagonia, Argentina. *J. South Am. Earth Sci.* **97**, 102401 (2020).
  46. Ibiricu, L. M. et al. New hadrosaurid (Dinosauria: Ornithomimidae) fossils from the uppermost Cretaceous of central Patagonia and the influence of paleoenvironment on South American hadrosaur distribution. *J. South Am. Earth Sci.* **110**, 103369 (2021).
  47. Casal, G. A., Allard, J. O. & Foix, N. Análisis estratigráfico y paleontológico del Cretácico Superior en la Cuenca del Golfo San Jorge: nueva unidad litoestratigráfica para el Grupo Chubut. *Rev. Asociación Geol. Argent.* **72**, 77–95 (2015).
  48. Tucker, R. T. et al. Age, depositional history, and paleoclimatic setting of Early Cretaceous dinosaur assemblages from the Sao Khua Formation (Khorat Group), Thailand. *Palaeogeogr. Palaeoclimatol. Palaeoecol.* **601**, 111107 (2022).
  49. Méndez, A. H., Novas, F. E. & Iori, F. V. First record of Megaraptora (Theropoda, Neovenatoridae) from Brazil. *Comptes Rendus Palevol* **11**, 251–256 (2012).
  50. Martinelli, A. G. et al. Insight on the theropod fauna from the Uberaba Formation (Bauru Group), Minas Gerais State: new megaraptoran specimen from the Late Cretaceous of Brazil. *Riv. Ital. di Paleontol. Stratigrafia* **119**, 205–214 (2013).

51. Sales, M. A. F. et al. New dinosaur remains and the tetrapod fauna from the Upper Cretaceous of Mato Grosso State, central Brazil. *Historical Biol.* **30**, 661–676 (2018).
52. Davis, S. N. et al. New records of Theropoda from a Late Cretaceous (Campanian–Maastrichtian) locality in the Magallanes–Austral Basin, Patagonia, and insights into end Cretaceous theropod diversity. *J. South Am. Earth Sci.* **122**, 104163 (2023).
53. Agnolín, F. L. et al. Fossil vertebrates from the Cerro Fortaleza Formation (Upper Cretaceous), Santa Cruz Province, Argentina. *Cretac. Res.* **154**, 105735 (2024).
54. Paulina-Carabajal, A., Barrios, F. T., Méndez, A. H., Cerda, I. A. & Lee, Y.-N. A Late Cretaceous dinosaur and crocodyliform faunal association–based on isolate teeth and osteoderms–at Cerro Fortaleza Formation (Campanian–Maastrichtian) type locality, Santa Cruz, Argentina. *PLoS ONE* **16**, e256233 (2021).
55. Casal, G. A., Martínez, R. D., Luna, M. & Ibiricu, L. M. Ordenamiento y caracterización faunística del Cretácico Superior del Grupo Chubut, Cuenca del Golfo San Jorge, Argentina. *Rev. Brasileira Paleontol.* **19**, 53–70 (2016).
56. Brusatte, S. L., Carr, T. D. & Norell, M. A. The osteology of *Alioramus*, a gracile and long-snouted tyrannosaurid (Dinosauria: Theropoda) from the Late Cretaceous of Mongolia. *Bull. Am. Mus. Nat. Hist.* **366**, 1–197 (2012).
57. Novas, F. E., Pol, D., Canale, J. I., Porfiri, J. D. & Calvo, J. O. A bizarre Cretaceous theropod dinosaur from Patagonia and the evolution of Gondwanan dromaeosaurids. *Proc. R. Soc. B: Biol. Sci.* **276**, 1101–1107 (2008).
58. Hendrickx, C., Araújo, R. & Mateus, O. The non-avian theropod quadrate I: standardized terminology with an overview of the anatomy and function. *PeerJ* **3**, e1245 (2015).
59. White, M. A., Bell, P. R., Cook, A. G., Poropat, S. F. & Elliott, D. A. The dentary of *Australovenator wintonensis* (Theropoda, Megaraptoridae); implications for megaraptorid dentition. *PeerJ* **3**, e1512 (2015).
60. Birch, S. A., Smith, E. T. & Bell, P. R. New theropod (Tetanurae: Avetheropoda) material from the ‘mid’-Cretaceous Griman Creek Formation at Lightning Ridge, New South Wales, Australia. *R. Soc. Open Sci.* **6**, 180826 (2019).
61. White, M. A. et al. New theropod remains and implications for megaraptorid diversity in the Winton Formation (lower Upper Cretaceous), Queensland, Australia. *R. Soc. Open Sci.* **7**, 191462 (2020).
62. Aranciaga Rolando, A. M., García Marsà, J. & Novas, F. Histology and pneumaticity of *Aoniraptor libertatem* (Dinosauria, Theropoda), an enigmatic mid-sized megaraptoran from Patagonia. *J. Anat.* **237**, 741–756 (2020).
63. Apesteguía, S., Smith, N. D., Juárez Valieri, R. & Makovicky, P. J. An unusual new theropod with a didactyl manus from the Upper Cretaceous of Patagonia, Argentina. *PLoS ONE* **11**, e0157793 (2016).
64. Calvo, J. O. et al. Morphological and phylogenetic significance of the first adult humerus of the Patagonian Cretaceous theropod *Megaraptor namunhualiquii* Novas, 1998. *Ann. Carnegie Mus.* **90**, 161–181 (2025).
65. White, M. A. et al. New forearm elements discovered of holotype specimen *Australovenator wintonensis* from Winton, Queensland, Australia. *PLoS ONE* **7**, e39364 (2012).
66. Rauhut, O. W. M. The interrelationships and evolution of basal theropod dinosaurs. *Spec. Pap. Palaeontol.* **69**, 1–213 (2003).
67. Chinsamy-Turan, A. *The Microstructure of Dinosaur Bone: Deciphering Biology Through Fine Scale Techniques* (The Johns Hopkins University Press, 2005).
68. Erickson, G. M. On dinosaur growth. *Annu. Rev. Earth Planet. Sci.* **42**, 675–697 (2014).
69. de Buffrénil, V. & Quilhac, A. Bone tissue types: a brief account of currently used categories. Pp. 147–190, in *Comparative Skeletal Histology and Palaeohistology* (V. de Buffrénil, L. Zylberberg, K. Padian, and A. de Ricqlès, eds.). (CRC Press, 2021).
70. Horner, J. R., de Ricqlès, A. & Padian, K. Variation in dinosaur skeletochronology indicators: implications for age assessment and physiology. *Paleobiology* **25**, 295–304 (1999).
71. Woodward, H. N., Horner, J. R. & Farlow, J. O. Osteohistological evidence for determinate growth in the American alligator. *J. Herpetol.* **45**, 339–342 (2011).
72. Sander, P. M. Long bone histology of the Tendaguru sauropods: implications for growth and biology. *Paleobiology* **26**, 466–488 (2000).
73. Erickson, G. M., Curry Rogers, K. A., Varricchio, D. J., Norell, M. A. & Xu, X. Growth patterns in brooding dinosaurs reveals the timing of sexual maturity in non-avian dinosaurs and genesis of the avian condition. *Biol. Lett.* **3**, 558–561 (2007).
74. Lee, A. H. & Werning, S. Sexual maturity in growing dinosaurs does not fit reptilian growth models. *Proc. Natl. Acad. Sci. USA* **105**, 582–587 (2008).
75. Castanet, J. & Baez, M. Adaptation and evolution in *Gallotia* lizards from the Canary Islands: age, growth, maturity and longevity. *Amphib. Reptilia* **12**, 81–102 (1991).
76. Khonsue, W., Chaiananporn, T. & Pomchote, P. Skeletochronological assessment of age in the Himalayan crocodile newt, *Tylotriton verrucosus* (Anderson, 1871) from Thailand. *Tropical Nat. Hist.* **10**, 181–188 (2010).
77. Xu, X., Tan, Q., Wang, J., Zhao, X. & Tan, L. A gigantic bird-like dinosaur from the Late Cretaceous of China. *Nature* **447**, 844–847 (2007).
78. Novas, F. E., Brissón Egli, F., Agnolín, F., Gianechini, F. & Cerda, I. A. Postcranial osteology of a new specimen of *Buitreraptor gonzalezorum* (Theropoda, Unenlagiidae). *Cretac. Res.* **83**, 127–167 (2018).
79. Canale, J. I., Cerda, I. A., Novas, F. E. & Haluza, A. Small-sized abelisaurid (Theropoda: Ceratosauria) remains from the Upper Cretaceous of northwest Patagonia, Argentina. *Cretac. Res.* **62**, 18–28 (2016).
80. Waskow, K. & Mateus, O. Dorsal rib histology of dinosaurs and a crocodylomorph from western Portugal: skeletochronological implications on age determination and life history traits. *Comptes Rendus Palevol* **16**, 425–439 (2017).
81. Waskow, K. & Sander, P. M. Growth record and histological variation in the dorsal ribs of *Camarasaurus* sp. (Sauropoda). *J. Vertebrate Paleontol.* **34**, 852–869 (2014).
82. Woodruff, D. C. & Foster, J. R. The first specimen of *Camarasaurus* (Dinosauria: Sauropoda) from Montana: the northernmost occurrence of the genus. *PLoS ONE* **12**, e0177423 (2017).
83. Woodward, H. N. & Lehman, T. M. Bone histology and micro-anatomy of *Alamosaurus sanjuanensis* (Sauropoda: Titanosauria) from the Maastrichtian of Big Bend National Park, Texas. *J. Vertebrate Paleontol.* **29**, 807–821 (2009).
84. Méndez, A. H. et al. New furileusaurian remains from La Invernada (northern Patagonia): a site of unusual abelisaurids abundance. *Cretac. Res.* **129**, 104989 (2022).
85. Brum, A. S., Bandeira, K. L. N., Sayão, J. M., Campos, D. A. & Kellner, A. W. A. Microstructure of axial bones of lithostrotian titanosaurs (Neosauropoda: Sauropodomorpha) shows extended fast-growing phase. *Cretac. Res.* **136**, 105220 (2022).
86. Wedel, M. J. Vertebral pneumaticity, air sacs, and the physiology of sauropod dinosaurs. *Paleobiology* **29**, 243–255 (2003).
87. O’Connor, P. M. Postcranial pneumaticity: an evaluation of soft-tissue influences on the postcranial skeleton and the reconstruction of pulmonary anatomy in archosaurs. *J. Morphol.* **267**, 1199–1226 (2006).

88. Claessens, L. P. A. M., O'Connor, P. M. & Unwin, D. M. Respiratory evolution facilitated the origin of pterosaur flight and aerial gigantism. *PLoS ONE* **4**, e4497 (2009).
89. Gianechini, F. A. & Zurriaguz, V. L. Vertebral pneumaticity of the paravian theropod *Unenlagia comahuensis*, from the Upper Cretaceous of Patagonia, Argentina. *Cretac. Res.* **127**, 104925 (2021).
90. Lambert, M., Bertozzo, F. & Sander, P. M. Bone histological correlates for air sacs and their implications for understanding the origin of the dinosaurian respiratory system. *Biol. Lett.* **14**, 20170514 (2018).
91. Chinsamy, A., Cerda, I. A. & Powell, J. Vascularised endosteal bone tissue in armoured sauropod dinosaurs. *Sci. Rep.* **6**, 24858 (2016).
92. Martínez, R. D. F. et al. A basal lithostrotian titanosaur (Dinosauria: Sauropoda) with a complete skull: implications for the evolution and paleobiology of Titanosauria. *PLoS ONE* **11**, e0151661 (2016).
93. Lamanna, M. C., Casal, G. A., Ibiricu, L. M. & Martínez, R. D. F. Non-avian dinosaur faunas of the Upper Cretaceous Bajo Barreal and Lago Colhué Huapi formations of central Patagonia, Argentina: paleobiogeographic and biostratigraphic implications. *Actas XXI Congreso Geológico Argentino*, 259–260 (2022).
94. Reguero, M. A. & Goin, F. J. Paleogeography and biogeography of the Gondwanan final breakup and its terrestrial vertebrates: new insights from southern South America and the “double Noah’s Ark” Antarctic Peninsula. *J. South Am. Earth Sci.* **108**, 103358 (2021).
95. Vajda, V. & Bercovici, A. The global vegetation pattern across the Cretaceous–Paleogene mass extinction interval: a template for other extinction events. *Glob. Planet. Change* **122**, 29–49 (2014).
96. Drumheller, S. K., McHugh, J. B., Kane, M., Riedel, A. & D’Amore, D. C. High frequencies of theropod bite marks provide evidence for feeding, scavenging, and possible cannibalism in a stressed Late Jurassic ecosystem. *PLoS ONE* **15**, e0233115 (2020).
97. White, M. A. et al. Forearm range of motion in *Australovenator wintonensis* (Theropoda, Megaraptoridae). *PLoS ONE* **10**, e0137709 (2015).
98. Yun, C.-G. Evaluating the paleoecology of the Megaraptora (Dinosauria: Theropoda) through biomechanical approaches. *Span. J. Palaeontol.* **39**, 223–234 (2024).
99. Burch, S. H. Complete forelimb myology of the basal theropod dinosaur *Tawa hallae* based on a novel robust muscle reconstruction method. *J. Anat.* **225**, 271–297 (2014).
100. Burch, S. H. Myology of the forelimb *Majungasaurus crenatissimus* (Theropoda, Abelisauridae) and the morphological consequences of extreme limb reduction. *J. Anat.* **231**, 515–531 (2017).
101. Porfiri, J. D. et al. *Diuqin lechiguanae* gen. et sp. nov., a new unenlagiine (Theropoda: Paraves) from the Bajo de la Carpá Formation (Neuquén Group, Upper Cretaceous) of Neuquén Province, Patagonia, Argentina. *BMC Ecol. Evol.* **24**, 77 (2024).
102. Campione, N. E. & Evans, D. C. A universal scaling relationship between body mass and proximal limb bone dimensions in quadrupedal terrestrial tetrapods. *BMC Biol.* **10**, 1–22 (2012).
103. Benson, R. B. J. et al. Rates of dinosaur body mass evolution indicate 170 million years of sustained ecological innovation on the avian stem lineage. *PLoS Biol.* **12**, e1001853 (2014).
104. Benson, R. B. J., Rich, T. H., Vickers-Rich, P. & Hall, M. Theropod fauna from southern Australia indicates high polar diversity and climate-driven dinosaur provinciality. *PLoS ONE* **7**, e37122 (2012).
105. White, M. A. & Campione, N. E. A three-dimensional approach to visualize pairwise morphological variation and its application to fragmentary palaeontological specimens. *PeerJ* **9**, e10545 (2021).
106. Morrison, C. et al. Rise of the king: Gondwanan origins and evolution of megaraptoran dinosaurs. *R. Soc. Open Sci.* **12**, 242238 (2025).
107. Chinsamy-Turan, A. & Raath, M. A. Preparation of fossil bone for histological examination. *Palaeontol. Afr.* **29**, 39–44 (1992).
108. Francillon-Vieillot, H. et al. Microstructure and mineralization of vertebrate skeletal tissues. In *Skeletal Biomineralization: Patterns, Processes and Evolutionary Trends* (Vol. 1) (ed. J. G. Carter). (Van Nostrand Reinhold, 1990).
109. Goloboff, P. A., Farris, J. S. & Nixon, K. C. TNT: a free program for phylogenetic analysis. *Cladistics* **24**, 774–786 (2008).
110. Walker, J. D. & Geissman, J. W. (compilers). *Geologic Time Scale v. 6.0*. Geological Society of America, Boulder. <https://doi.org/10.1130/2022.CTS006C> (2022).

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## Author contributions

L.M.I. contributed to the excavation of the *Joaquinraptor casali* quarry in the field, designed and led all aspects of the study (with the exception of the histological analysis), wrote the initial drafts of nearly all sections of the manuscript, and contributed to the editing of subsequent drafts. M.C.L. contributed to most aspects of the study (e.g., field excavation, osteological description, phylogenetic analysis, palaeobiogeographic and palaeobiological observations), extensively edited and contributed to multiple drafts of the manuscript, and supervised the construction of most figures. B.N.A. discovered the holotype of *J. casali* and contributed to its excavation in the field and to the geological aspects of the study. I.A.C. designed and led the histological analysis and wrote the initial draft of the corresponding section of the manuscript. J.L.C. and N.V.C. contributed in the field and formatted the final version of the manuscript. M.L. contributed in the field and mechanically prepared and photographed all fossils. R.D.M. reviewed the initial draft and the originally submitted version of the manuscript prior to his passing.

## Competing interests

The authors declare no competing interests.



## Additional information

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